Inventor Search

Fubara 10/071,490

13/04/2006

=> d ibib abs 11 1-2

L1 ANSWER 1 OF 2 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2003:609852 HCAPLUS

DOCUMENT NUMBER: 139:154974

TITLE: Compositions and methods for forming and strengthening

bone

INVENTOR(S): Marchosky, J. Alexander

PATENT ASSIGNEE(S): USA

SOURCE: U.S. Pat. Appl. Publ., 16 pp.

CODEN: USXXCO

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
				
US 2003147860	A1	20030807	US 2002-71490	20020207
PRIORITY APPLN. INFO.:			US 2002-71490	20020207

AB Compns. are provided which stimulate bone growth. Also provided are methods for utilizing the compns. for filling in bone defects, promoting rapid fusion of bone fractures, grafts, and bone-prostheses, and promoting strengthening of osteoporotic bones. The appearance of bone formation at the site of bone defect in rat's femur was shown after application of a composition containing demineralized bone matrix, hyaluronic acid, and purified vascular endothelial growth factor at 12 wk.

L1 ANSWER 2 OF 2 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2001:12589 HCAPLUS

DOCUMENT NUMBER:

134:76442

TITLE:

Compositions containing growth factors and methods for

forming and strengthening bone

INVENTOR(S): Marchosky, J. Alexander

PATENT ASSIGNEE(S): USA

SOURCE: PCT Int. Appl., 41 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PAT	CENT	NO.			KIN	D	DATE			APPL	ICAT	ION	NO.		Di	ATE	
WO	2001	0007	92		A1	-	2001	0104		WO 2	000-	US17	955		2	0000	629
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		HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	ΚZ,	LC,	LK,	LR,	LS,	LT,
		LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	NZ,	PL,	PT,	RO,	RU,
		SD,	SE,	SG,	SI,	SK,	SL,	ТJ,	TM,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VN,
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		DE,	DK,	ES,	FI,	FR,	GB,	GR,	IE,	ΙT,	LU,	MC,	NL,	PT,	SE,	BF,	ВJ,
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CA	2377	435			AΑ		2001	0104		CA 2	000-	2377	435		2	0000	629
US	6372															0000	
EΡ	1203	074			A1		2002	0508		EP 2	-000	9433	09		20	0000	629
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		ΙE,	SI,	LT,	LV,	FI,	RO,	MK,	CY,	AL							
ΑU	7823	94			В2		2005	0721		AU 2	000-	5779	9		2	0000	629

13/04/2006

Fubara 10/071,490

PRIORITY APPLN. INFO.:

US 1999-141386P P 19990629 WO 2000-US17955 W 20000629

AB Compns. for stimulating bone growth comprise (a) growth factors, (b) demineralized, non-decalcified bone matrix, (c) a scaffolding material selected from cancelous bone, chitosan, chitosan-protein, and chitosan-protein fibers, and (d) a gel material selected from chitosan and its derivs., alginate, or hyaluronic acid. Addnl., compns. may contain angiogenesis-stimulating materials and osteoinductive materials. Methods for utilizing the compns. for filling in bone defects, promoting rapid fusion of bone fractures, grafts, and bone-prostheses, and promoting strengthening of osteoporotic bones are also provided. For example, bone formation at the site of bone defect was observed 12 wk after the application of the composition containing demineralized bone matrix, hyaluronic acid, and vascular endothelial growth factor.

REFERENCE COUNT:

THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

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(FILE 'HOME' ENTERED AT 17:21:18 ON 13 APR 2006)

FILE 'HCAPLUS' ENTERED AT 17:22:38 ON 13 APR 2006

E MARCHOSKY J ALEX/AU

2 SEA ABB=ON "MARCHOSKY J ALEXANDER"/AU L1ANALYZE L1 1-2 CT : 13 TERMS L2

FILE 'REGISTRY' ENTERED AT 17:25:36 ON 13 APR 2006

E HYALURONIC ACID/CN

1 SEA ABB=ON "HYALURONIC ACID"/CN L3

FILE 'HCAPLUS' ENTERED AT 17:25:54 ON 13 APR 2006 16360 SEA ABB=ON L3 OR ?HYALURONIC?(W)?ACID?

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10 SEA ABB=ON L5 AND ?ALLOGRAFT? L6

23 SEA ABB=ON L4 AND (?BONE?(3A)?FORM?)(5A)?INDUC? L7

33 SEA ABB=ON L6 OR L7 $\Gamma8$

L9 20 SEA ABB=ON L8 AND (?GROW? OR ?STRENGTH?)

L10 33 SEA ABB=ON L8 OR L9

16 SEA ABB=ON L10 AND (PRD<19990629 OR PD<19990629)

// Le cife from CAPlus L11

JICST-EPLUS' ENTERED AT 17:30:50 ON FILE 'MEDLINE, BIOSIS, EMBASE, JAPIO,

13 APR 2006

L12 12 SEA ABB=ON L11 6 DUP REMOV L12 (6 DUPLICATES REMOVED) 6 citation L13

FILE 'USPATFULL' ENTERED AT 17:34:33 ON 13 APR 2006

215 SEA ABB=ON L10 AND (PRD<19990629 OR PD<19990629)

L14

215 SEA ABB=ON L14 AND (?GROW? OR ?STRENGTH?) L15

40 SEA ABB=ON L15 AND ?ALLOGRAFT?
40 SEA ABB=ON L16 AND ?FORM? FORM? FORM? GETTER HEAD (15 Part of the season of t L16 L17

FILE HOME

FILE HCAPLUS

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FILE COVERS 1907 - 13 Apr 2006 VOL 144 ISS 16 FILE LAST UPDATED: 12 Apr 2006 (20060412/ED)

New CAS Information Use Policies, enter HELP USAGETERMS for details.

This file contains CAS Registry Numbers for easy and accurate substance identification.

FILE REGISTRY

Property values tagged with IC are from the ZIC/VINITI data file

provided by InfoChem.

STRUCTURE FILE UPDATES: 11 APR 2006 HIGHEST RN 880129-32-8 DICTIONARY FILE UPDATES: 11 APR 2006 HIGHEST RN 880129-32-8

New CAS Information Use Policies, enter HELP USAGETERMS for details.

TSCA INFORMATION NOW CURRENT THROUGH January 6, 2006

Please note that search-term pricing does apply when conducting SmartSELECT searches.

Structure search iteration limits have been increased. See HELP SLIMITS for details.

REGISTRY includes numerically searchable data for experimental and predicted properties as well as tags indicating availability of experimental property data in the original document. For information on property searching in REGISTRY, refer to:

http://www.cas.org/ONLINE/UG/regprops.html

FILE MEDLINE

FILE LAST UPDATED: 12 APR 2006 (20060412/UP). FILE COVERS 1950 TO DATE.

On December 11, 2005, the 2006 MeSH terms were loaded.

The MEDLINE reload for 2006 is now (26 Feb.) available. For details on the 2006 reload, enter HELP RLOAD at an arrow prompt (=>). See also:

http://www.nlm.nih.gov/mesh/

http://www.nlm.nih.gov/pubs/techbull/nd04/nd04_mesh.html

http://www.nlm.nih.gov/pubs/techbull/nd05/nd05 med data changes.html

http://www.nlm.nih.gov/pubs/techbull/nd05/nd05 2006 MeSH.html

OLDMEDLINE is covered back to 1950.

MEDLINE thesauri in the /CN, /CT, and /MN fields incorporate the MeSH 2006 vocabulary.

This file contains CAS Registry Numbers for easy and accurate substance identification.

FILE BIOSIS

FILE COVERS 1969 TO DATE.

CAS REGISTRY NUMBERS AND CHEMICAL NAMES (CNs) PRESENT FROM JANUARY 1969 TO DATE.

RECORDS LAST ADDED: 12 April 2006 (20060412/ED)

FILE EMBASE

FILE COVERS 1974 TO 13 Apr 2006 (20060413/ED)

EMBASE has been reloaded. Enter HELP RLOAD for details.

EMBASE is now updated daily. SDI frequency remains weekly (default) and biweekly.

This file contains CAS Registry Numbers for easy and accurate substance identification.

FILE JAPIO

FILE LAST UPDATED: 3 APR 2006 <20060403/UP>
FILE COVERS APRIL 1973 TO DECEMBER 22, 2005

- >>> GRAPHIC IMAGES AVAILABLE <<<
- >>> NEW IPC8 DATA AND FUNCTIONALITY NOT YET AVAILABLE IN THIS FILE.

 USE IPC7 FORMAT FOR SEARCHING THE IPC. WATCH THIS SPACE FOR FURTHER

 DEVELOPMENTS AND SEE OUR NEWS SECTION FOR FURTHER INFORMATION

 ABOUT THE IPC REFORM <<<

FILE JICST-EPLUS FILE COVERS 1985 TO 11 APR 2006 (20060411/ED)

THE JICST-EPLUS FILE HAS BEEN RELOADED TO REFLECT THE 1999 CONTROLLED TERM (/CT) THESAURUS RELOAD.

FILE USPATFULL

FILE COVERS 1971 TO PATENT PUBLICATION DATE: 13 Apr 2006 (20060413/PD)
FILE LAST UPDATED: 13 Apr 2006 (20060413/ED)
HIGHEST GRANTED PATENT NUMBER: US7028340
HIGHEST APPLICATION PUBLICATION NUMBER: US2006080750
CA INDEXING IS CURRENT THROUGH 13 Apr 2006 (20060413/UPCA)
ISSUE CLASS FIELDS (/INCL) CURRENT THROUGH: 13 Apr 2006 (20060413/PD)
REVISED CLASS FIELDS (/NCL) LAST RELOADED: Feb 2006
USPTO MANUAL OF CLASSIFICATIONS THESAURUS ISSUE DATE: Feb 2006

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L7
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L11
=> d ibib abs 111 1-16
L11 ANSWER 1 OF 16 HCAPLUS COPYRIGHT 2006 ACS on STN
                         2006:138967 HCAPLUS
ACCESSION NUMBER:
                         144:199025
DOCUMENT NUMBER:
TITLE:
                        Demineralized corticocancellous
                        bone sheet
```

INVENTOR(S):

Sunwoo, Moon Hae; Gertzman, Arthur A.; Stroever, Bruce

PATENT ASSIGNEE(S):

Musculoskeletal Transplant Foundation, USA

U.S., 9 pp., Cont.-in-part of U.S. Ser. No. 413,815. SOURCE:

CODEN: USXXAM

DOCUMENT TYPE: Patent English LANGUAGE: FAMILY ACC. NUM. COUNT: 10

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 6998135	В1	20060214	US 2001- § 53761	20010514 <
US 6030635	Α.	20000229	US 1998-31750	19980227
US 6326018	B1	20011204	US 1999-413815	19991007 <
EP 1477176	A1	20041117	EP 2004-77080	20000222
R: AT, BE, CH,	DE, DK	, ES, FR, GB	, GR, IT, LI, LU, NL,	SE, MC, PT,
IE, FI, CY				
AT 297766	E	20050715	AT 2000-301370	20000222
ES 2241549	Т3	20051101	ES 2000-301370	20000222
PRIORITY APPLN. INFO.:			US 1998-31750 A	1 19980227 <
			US 1999-413815 A	2 19991007
			EP 2000-301370 A	3 20000222

AB A flexible, demineralized unitary bone sheet comprised of cortical cancellous bone having a residual calcium weight of 3.0% to 8.0% with a hyaluronic acid component

having a mol. weight of 700,000 to 1,500,000 with the weight of the same ranging

from 1% to about 5% of the total sheet weight. The bone sheet is adapted for use during the in vivo repair of a mammalian or animal skeletal system with the thickness of the cortical cancellous sheet ranging from 2.0 mm to about 8.0 mm. The bone sheet has sufficient flexibility to allow the sheet to be shaped to conform to the configuration of a skeletal region to be repaired and sufficient tensile strength to allow the sheet to be so shaped without damage to the sheet. Preparation of cortical strips from human femoral allograft tissue taken from a qualified donor is described.

REFERENCE COUNT:

17 THERE ARE 17 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L11 ANSWER 2 OF 16 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

2004:402760 HCAPLUS

DOCUMENT NUMBER:

140:380720

TITLE:

Synthetic bone matrix implant comprising osteogenic protein 1 in a crosslinked collagen-glycosaminoglycan

INVENTOR(S):

Kuberasampath, Thangavel; Tarrant, Lawrence Berlowitz

Stryker Corp., USA

PATENT ASSIGNEE(S): SOURCE:

U.S. Pat. Appl. Publ., 10 pp., Cont. of U.S. Ser. No.

104,865.

CODEN: USXXCO

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATE	ENT NO.	KIND	DATE	APPLICATION NO.	DATE
	2002151985	A1	20021017	US 2001-882875	20010615 <
	5605117 539574	B2 B2	20030812 19930729	AU 1991-79614	19910522 <
	06505642 508211	T2 A1	19940630 19940803	JP 1991-510269 EP 1991-911588	19910522 <
EP 6	508211	B1	19951227		an.
AT 1	R: AT, BE, CH, L32043	Ē	19960115	, GR, IT, LI, LU, NL, S AT 1991-911588	19910522 <
	2081484 2082946	T3 C	19960301 19961210	ES 1991-911588 CA 1991-2082946	19910522 < 19910522 <
PRIORITY	APPLN. INFO.:				3 19900529 < 1 19950518 <
				US 1998-104865 A	1 19980625 <
				WO 1991-US3603 A	19910522 <

Disclosed is an osteogenic device capable of inducing the AB formation of endochondral bone in a shape conforming substantially to the shape of the device when implanted in a mammalian host. The device includes an osteogenic protein (OP1) dispersed within a porous matrix comprising a polymer of collagen and glycosaminoglycan crosslinked to an Mc value of about 800 to about 60,000. Also disclosed are a method of inducing mammalian bone growth, and a method of inducing conductive bone growth from viable mammalian bone.

L11 ANSWER 3 OF 16 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2003:1004372 HCAPLUS

140:8875 DOCUMENT NUMBER:

TITLE:

INVENTOR(S):

Assembled implant including mixed-composition segment Bianchi, John R.; Mills, C. Randal; Gorham, P. J.;

Esch, Michael; Carter, Kevin C.; Coleman, Pat; Ross, Kevin; Rambo, Harry W.; Jones, Darren G.; Buskirk, Dayna; Donda, Russell S.

PATENT ASSIGNEE(S): USA

U.S. Pat. Appl. Publ., 36 pp., Cont.-in-part of U.S. SOURCE:

Appl. 2001 31,254.

CODEN: USXXCO

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT: 11

PATENT INFORMATION:

APPLICATION NO. DATE PATENT NO. KIND DATE

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US 2001-941154
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    US 2002106393
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             HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS,
             LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO,
             RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, UZ, VN,
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                                                                   20010907
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     US 2004115172
                                                                P 20000210
                                            US 2000-181622P
PRIORITY APPLN. INFO.:
                                                                A2 20010212
                                            US 2001-782594
                                            US 1998-191132
                                                                A2 19981113 <--
                                                                A2 19981113 <--
                                            US 1998-191232
                                                                B1 19990728
                                            US 1999-363909
                                                                A1 19990809
                                            US 1999-370194
                                            US 1999-378527
                                                                A2 19990820
                                            US 2000-481319
                                                                A1 20000111
                                                                A1 20000317
                                            US 2000-528034
                                                                A1 20000512
                                            US 2000-DS123227
                                            US 2001-941154
                                                                A 20010827
                                                                W 20010907
                                            WO 2001-US27683
     This invention provides a method for manufacture of autograft,
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AΒ allograft and xenograft implants which comprises assembling such implants from smaller pieces of graft materials to form a larger graft implant product. One segment of an assembled graft implant is comprised of 2 or more discrete regions containing at least one synthetic segment and at least one demineralized bone segment and having distinct characteristics and/or properties. The synthetic segment is comprised of e.g., stainless steel, titanium, nylon, polycarbonate, polypropylene, polyacetal, PEG, polyvinylpyrolidone, polyacrylates, polyesters, and polysulfone's.

L11 ANSWER 4 OF 16 HCAPLUS COPYRIGHT 2006 ACS on STN

2003:874783 HCAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 139:354539

TITLE:

Malleable putty and flowable paste with allograft bone having residual calcium for

filling bone defects

Gertzman, Arthur A.; Sunwoo, Moon H. INVENTOR(S):

Musculoskeletal Transplant Foundation, USA PATENT ASSIGNEE(S): SOURCE:

U.S. Pat. Appl. Publ., 16 pp., Cont.-in-part of U.S.

6,437,018. CODEN: USXXCO

Patent DOCUMENT TYPE: English LANGUAGE:

10 FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 2003206937	A1	20031106	US 2001-983526	20011024 <

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US 6911212
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     EP 1477176
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PRIORITY APPLN. INFO.:
                                                                 B2 19990803
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                                                                 A2 20000229
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                                             EP 2000-301370
                                                                 A3 20000222
                                                                 A2 20011024
                                             US 2001-983526
AB
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AB The invention is directed toward a malleable bone putty and a flowable pastel composition for application to a bone defect site to promote new bone growth at the site which comprises a new bone

growth inducing compound of partially demineralized
lyophilized allograft bone material having a residual

calcium content of 4-8% dry weight. The bone powder has a particle size of $100\text{--}800~\mu$ and is mixed in a high mol. weight hydrogel carrier containing a sodium phosphate saline buffer, the hydrogel component of the carrier at 1.00--50% of the composition and having a mol. weight of about at least 700,000 Daltons. The composition has a pH of 6.8-7.4 contains 25-35% bone powder and can be addnl. provided with BMP's. A malleable putty of 4%

hyaluronic acid was prepared by mixing freeze dried

demineralized cortical allograft bone of

particle size of $100-800 \mu$ with 146.6 g of a 4% solution of

hyaluronic acid (mol. wt, 700,000 Daltons) in phosphate

buffered saline. The bone component was added to achieve a bone concentration of

approx. 32%. The solution was well mixed and allowed to stand for 2-3 h at room temperature. This provides a malleable putty with a penetration unit of 66 and excellent formability properties and a pH of 7.0.

REFERENCE COUNT: 13 THERE ARE 13 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L11 ANSWER 5 OF 16 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2002:748712 HCAPLUS

DOCUMENT NUMBER: 137:268501

TITLE: Malleable paste with allograft bone

reinforcement for filling bone defects Gertzman, Arthur A.; Sunwoo, Moon Hae

INVENTOR(S): Gertzman, Arthur A.; Sunwoo, Moon Hae PATENT ASSIGNEE(S): Musculoskeletal Transplant Foundation, USA

SOURCE: U.S., 11 pp., Cont.-in-part of U. S. Ser. No. 515,656.

CODEN: USXXAM

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 10

PATENT INFORMATION:

PATENT NO.	KIND DATE	APPLICATION NO.	DATE
US 6458375	B1 20021001	US 2000-677891	20001003 <
US 6030635	A 20000229	US 1998-31750	19980227
EP 1477176	A1 20041117	EP 2004-77080	20000222
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AT 297766	E 20050715	AT 2000-301370	20000222
ES 2241549	T3 20051101	ES 2000-301370	20000222

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US 6437018
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PRIORITY APPLN. INFO.:
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                                                                 A2 19980227 <--
                                                                 A2 20000229
                                             US 2000-515656
                                             US 1999-365880
                                                                B2 19990803
                                             EP 2000-301370
                                                                 A3 20000222
                                             US 2000-677891
                                                                 A2 20001003
                                             US 2000-739214
                                                                 A 20001219
                                             WO 2001-US27744
                                                                 W 20011002
                                             WO 2001-US27745
                                                                 W
                                                                    20011002
     The invention is directed toward a sterile malleable bone composition for
AΒ
     application to a bone defect site to promote new bone growth at
    the site comprising a mixture of demineralized osteogenic bone powder with a particle size ranging from about 250 to about
     750 \mu and surface demineralized cortical bone rods
    having a diameter ranging from 1.0 mm to 5.00 mm or larger bone chips. The
     surface demineralized cortical bone rods have diameter to
     length ratio ranging from 1:2 to 1:20. The demineralized
     bone powder range from about 25 to about 30% of the weight of the
     composition and the cortical bone rods range from 5% to about 10% of the
weight of
     the composition with the carrier being selected from the high mol. weight
     hydrogel, e.g., chitosan and sodium hyaluronate, in aqueous solution having a
     high mol. weight over 700,000 Dalton and ranging from about 2.0% to about
     5.0% by weight of the carrier solution For example, to a malleable putty of 4%
     solution of hyaluronic acid (mol. weight of 1,000,000
     Dalton) in phosphate buffered saline at a pH of about 7.2 and a 140,000
     viscosity containing 15-30% by weight of cortical allograft bone powder
     (particle size of 250-750 \mu) were added 5-25% mineralized bone chips
     (0.1-10 \text{ mm}) and mineralized rods (0.1-10 \text{ mm}) to obtain a paste.
                               THERE ARE 14 CITED REFERENCES AVAILABLE FOR THIS
REFERENCE COUNT:
                         14
                               RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
L11 ANSWER 6 OF 16 HCAPLUS COPYRIGHT 2006 ACS on STN
                         2002:637559 HCAPLUS
ACCESSION NUMBER:
                         137:175008
DOCUMENT NUMBER:
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Assembled implants prepared from mixed-composition segments made of natural bone, alloys, and plastics Bianchi, John R.; Mills, Randal C.; Gorham, P. J.;

Esch, Michael; Carter, Kevin C.; Coleman, Pat; Ross, Kevin; Rambo, Harry W.; Jones, Darren G.; Buskirk,

TITLE:

INVENTOR(S):

Dayna

PATENT ASSIGNEE(S): Regeneration Technologies, Inc., USA

SOURCE: PCT Int. Appl., 70 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 11

PATENT INFORMATION:

PA	TENT	NO.			KIN	D	DATE			APPL	ICAT	ION	NO.		D.	ATE	
WO	2002	0641	80		A1		2002	0822	,	WO 2	001-	US27	683		2	0010	907
	W:	ΑE,	AG,	AL,	AM,	AT,	ΑU,	AZ,	BA,	BB,	BG,	BR,	BY,	ΒZ,	CA,	CH,	CN,
		CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EE,	ES,	FI,	GB,	GD,	GE,	GH,	GM,
		•		•	•			JP,			•						
					•			MK,					-				
					SG,	SI,	SK,	SL,	ТJ,	TM,	TR,	TT,	TZ,	UA,	ÜG,	UZ,	VN,
			ZA,														
	RW:	GH,															
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US	2001	0312	54		A1												212 <
	2002							8080									
CA	2437	763			AA												
EP	1359	950			A1		2003	1112		EP 2	001-	9686	00		2	0010	907
	R:	ΑT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	ΙT,	LI,	LU,	NL,	SE,	MC,	PT,
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JP	2005	5102	58		Т2		2005	0421		JP 2	002-	5639	72		2	0010	907
PRIORIT	Y APP	LN.	INFO	.:						US 2	001-	7825	94		A 2	0010	212
										US 2	001-	9411	54		A 2	0010	827
										US 1	998-	1911	32		A2 1	9981	113 <
										US 2	000-	1816	22P		P 2	0000	210
										WO 2	001-	US27	683	1	W 2	0010	907

AB A method for manufacture of autograft, allograft and xenograft bone implants comprises assembling such implants from smaller pieces of bone graft materials to form a larger graft implant product. One segment of an assembled graft implant is comprised of two or more discrete regions having distinct characteristics and/or properties. An assembled graft implant comprises individual segments fastened together, the segments being mineralized bone, demineralized bone,

or a synthetic segment selected from alloys and plastic materials.

REFERENCE COUNT: 8 THERE ARE 8 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L11 ANSWER 7 OF 16 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2002:482945 HCAPLUS

DOCUMENT NUMBER: 137:52435

TITLE: Methods and articles for regenerating bone or

periodontal tissue

INVENTOR(S): White, Charles F.; Flynn, Charles; Cook, Alonzo D.;

Hardwick, William R.; Wikesjo, Ulf M. E.; Thomson,

Robert C.

PATENT ASSIGNEE(S): USA

SOURCE: U.S., 37 pp.

CODEN: USXXAM

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

DATE APPLICATION NO. DATE KIND PATENT NO. -------------------B1 20020625 US 1998-205150 19981203 <-- US 1998-205150 19981203 <--US 6409764 PRIORITY APPLN. INFO.: There are numerous medical situations involving deficiencies of living bone or periodontal tissue and where increase of living bone or periodontal tissue mass is desired. Methods are described wherein a configured, shell-like device that is capable of being penetrated by living cells and tissues, is implanted into the body of a mammal in such a way as to establish a space, the space being at least partly, bounded by the device. The configuration of the device is such that the configuration of the established space is essentially the same as the configuration of living bone or periodontal tissue that is desired for treatment of the tissue deficiency. At least one protein from the transforming growth factor- β (TGF- β) superfamily of proteins is placed within the established space for the purpose of stimulating the growth of living bone or periodontal tissue within the established space. A kit for the generation of living bone or periodontal tissue, comprised of the components mentioned above, is also disclosed. An example is given for periodontal regeneration with tissue exclusive and tissue penetrable devices containing TGF- β proteins and comprising PTFE devices. REFERENCE COUNT: THERE ARE 35 CITED REFERENCES AVAILABLE FOR THIS 35

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L11 ANSWER 8 OF 16 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2002:450194 HCAPLUS

DOCUMENT NUMBER: 1

137:24375

TITLE:

Bone graft substitute composition containing calcium

sulfate

INVENTOR(S):

Petersen, Donald W.; Richelsoph, Kelly Coupe; Haggard,

Warren Oliver; Hagan, Cary P.; Randolph, Donald A.

PATENT ASSIGNEE(S):

USA

SOURCE:

U.S. Pat. Appl. Publ., 8 pp., Cont.-in-part of U.S.

Ser. No. 327,761. CODEN: USXXCO

DOCUMENT TYPE: LANGUAGE: Patent English

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.		DATE
				_	
US 2002071827	A1	20020613	US 2001-915997		20010726 <
US 2002110541	A1	20020815	US 2002-60697		20020130 <
PRIORITY APPLN. INFO.:			US 1999-327761	A2	19990607 <
			US 2001-915997	A1	20010726

AB A bone graft substitute composition essentially made of calcium sulfate, a mixing solution, and a plasticizing substance, i.e., a cellulose derivative is described. A bone graft substitute composition can include demineralized bone matrix and cancellous bone. For example, an injectable bone graft substitute composition was prepared containing 100 parts by weight of medical grade calcium sulfate

hemihydrate, 11.1 parts by weight of CM-cellulose, 69.4 parts by weight of demineralized bone matrix, and 162 parts by weight of sterile water. The resultant injectable bone graft substitute composition was characterized by handability, ejectability, and robustness. The composition was well tolerated by the bone and healed a large medullary defect 30-100% at 6-20 wk with viable new bone in a canine bone defect model.

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L11 ANSWER 9 OF 16 HCAPLUS COPYRIGHT 2006 ACS on STN
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ACCESSION NUMBER: 2000:909060 HCAPLUS

DOCUMENT NUMBER: 134:61583

TITLE: Collagen matrix and growth factors in

non-immunogenic compositions for programming an organic matrix for remodeling into a target tissue

INVENTOR(S): Ashkar, Samy; Atala, Anthony

PATENT ASSIGNEE(S): Children's Medical Center Corp., USA

SOURCE: U.S., 10 pp., Cont.-in-part of U. S. Ser. No. 937,873.

CODEN: USXXAM

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 3

PATENT INFORMATION:

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KIND
                                DATE
                                       APPLICATION NO.
                                                                   DATE
     PATENT NO.
                                            _____
                                                                    _____
     _____
                         ____
                                _____
                                20001226 US 1998-58048
19980409 WO 1997-US17530
    US 6165487
                                                                    19980409 <--
                         Α
                                                                   19970929 <--
    WO 9814222
                         A1
            AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CZ, DE, DK,
             EE, ES, FI, GB, GE, GH, HU, ID, IL, IS, JP, KE, KG, KP, KR, KZ,
             LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL,
             PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, UZ,
             VN, YU, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
         RW: GH, KE, LS, MW, SD, SZ, UG, ZW, AT, BE, CH, DE, DK, ES, FI, FR,
             GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA,
             GN, ML, MR, NE, SN, TD, TG
                                            WO 1999-US7742
                                19991021
                                                                    19990408 <--
    WO 9952572
                          Α1
             AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ,
             DE, DK, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK,
             MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ,
             TM, TR, TT, UA, UG, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD,
             RU, TJ, TM
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             ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG,
             CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
     AU 9933875
                          A1
                                19991101
                                            AU 1999-33875
                                                                    19990408 <--
                                                               P 19960930 <--
PRIORITY APPLN. INFO.:
                                            US 1996-27123P
                                                                A2 19970929 <--
                                            US 1997-937873
                                                                A1 19970929 <--
                                            WO 1997-US17530
                                            US 1998-58048
                                                                A 19980409 <--
                                            WO 1999-US7742
                                                                 W 19990408 <--
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AB Methods for programming a non-immunogenic matrix for remodeling into a target tissue are disclosed. Also disclosed are compns. containing demineralized collagen and a growth factor, e.g., osteopontin, which can promote the growth of selected tissue types in a subject. Methods for preparing the compns. are also described. The methods and compns. are useful for treatment of defects in tissues such as bone, cartilage, and muscle. For example, a bone-forming matrix was prepared by suspending demineralized bone in a physiol. saline solution with 0.1% osteopontin, 0.01% bone sialoprotein, and 0.1% of high-mol.-weight hyaluronic acid and drying. The bone-forming matrix provided new bone formation in bone defects. It is believed that the bone forming compns. of the invention provided results equal to, or superior to, the results seen with bone allograft treatment.

REFERENCE COUNT: 22 THERE ARE 22 CITED REFERENCE

THERE ARE 22 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L11 ANSWER 10 OF 16 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2000:522637 HCAPLUS

DOCUMENT NUMBER: 133:109933

TITLE: Bone induction accelerator for treating fracture or

bone deficiency

INVENTOR(S): Kawachi, Toshiyuki; Takahashi, Makoto; Shinomiya,

Kenichi

PATENT ASSIGNEE(S): Tokyo Medical and Dental University, Japan

SOURCE: Jpn. Tokkyo Koho, 7 pp.

. CODEN: JTXXFF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 3032824	В1	20000417	JP 1999-19712	19990128
JP 2000212204	A2	20000802		
EP 1044691	A2	20001018	EP 2000-250026	20000127 <
EP 1044691	A3	20030611		
R: AT, BE, Ch	, DE, DE	K, ES, FR, GB	, GR, IT, LI, LU, N	L, SE, MC, PT,
IE, SI, LT	, LV, FI	[, RO		
US 2003064960	A1	20030403	US 2002-291658	20021112 <
PRIORITY APPLN. INFO.:			JP 1999-19712	A 19990128 <
•			US 2000-492704	B3 20000127

AB Bone induction accelerator effective in treating fracture or bone deficiency contains glycosaminoglycan-lipid conjugates or their pharmaceutically acceptable salts. Glycosaminoglycan is selected from hyaluronic acid, chondroitin, chodroitinsulfate, chondroitin polysulfate, dermatan sulfate, heparin, keratin sulfate and keratin polysulfate and lipid is phosphatidylethanolamine.

L11 ANSWER 11 OF 16 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2000:141482 HCAPLUS

DOCUMENT NUMBER: 132:185482

TITLE: Malleable paste for filling bone defects
INVENTOR(S): Gertzman, Arthur A.; Sunwoo, Moon Hae
PATENT ASSIGNEE(S): Musculoskeletal Transplant Foundation, USA

SOURCE: U.S., 7 pp.
CODEN: USXXAM

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 10

PATENT INFORMATION:

PATENT NO.	KIND DATE	APPLICATION NO.	DATE
US 6030635 US 6326018	A 20000229 B1 20011204	US 1998-31750 US 1999-413815	19980227 19991007 <
CA 2294686 CA 2294686	AA 20010706 C 20050503	CA 2000-2294686	20000106 <
EP 1127581	A1 20010829	EP 2000-301370	20000222 <
EP 1127581 R: AT, BE, CH,	B1 20050615 DE, DK, ES, FR,	GB, GR, IT, LI, LU, NL,	SE, MC, PT,
IE, SI, LT, EP 1477176	LV, FI, RO A1 20041117	EP 2004-77080	20000222
R: AT, BE, CH,		GB, GR, IT, LI, LU, NL,	
IE, FI, CY AT 297766	E 20050715	AT 2000-301370	20000222

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ES 2000-301370
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    ES 2241549
                          Т3
                                 20051101
    US 6437018
                                 20020820
                                             US 2000-515656
                                                                     20000229 <--
                          В1
    US 6458375
                                 20021001
                                             US 2000-677891
                                                                     20001003 <--
                          В1
    US 6998135
                          В1
                                 20060214
                                             US 2001-853761
                                                                     20010514 <--
                                20031106
                                             US 2001-983526
                                                                     20011024 <--
    US 2003206937
                         A1
                         B2
                                20050628
    US 6911212
                                             US 2002-84090
                                                                     20020228 <--
                         E
                                20040525
    US 38522
                                             US 2002-150097
                                                                     20020520 <--
    US 2002192263
                         A1
                                20021219
                         A1
                                20021226
                                             US 2002-222807
                                                                     20020819 <--
    US 2002197242
    US 7019192
                         B2
                                20060328
                                                             20040421 \
A3 19980227 <--
B2 19990803
A2 19991007
A3 20000222
                                                                     20040421 <--
     US 2004197373
                         A1
                                20041007
                                             US 2004-828316
                                             US 1998-31750
PRIORITY APPLN. INFO.:
                                             US 1999-365880
                                             US 1999-413815
                                             EP 2000-301370
                                             US 2000-515656
                                                                 A2 20000229
                                                                 A2 20011024
                                             US 2001-983526
                                             US 2002-222807
                                                                 A1 20020819
    The invention is directed toward a malleable bone putty and a flowable gel
AΒ
     composition for application to a bone defect site to promote new bone
    growth at the site which comprises a new bone
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growth inducing compound of demineralized lyophilized allograft bone powder. The bone powder has a particle size ranging from about 100 to about 850 μ and is mixed in a high mol.

weight hydrogel carrier, the hydrogel component of the carrier ranging from 0.3 to 3.0% of the composition and having a mol. weight of about at least 10,000

Daltons. The composition contains about 25% to about 40% bone powder and can be addnl. provided with BMP's and a sodium phosphate buffer. A malleable putty of 2% solution hyaluronic acid in isotonic saline with 250-420 μ cortical allograft bone powder at 30%. Freeze dried cortical allograft bone (502 mg) of particle size ranging $250-420~\mu$ was mixed into 1170 mg of a 2% solution of sodium hyaluronate in isotonic saline. The bone component is added to achieve a bone concentration

of 30% (weight/weight). The solution was well mixed and allowed to stand for 2-3

room temperature to provide a malleable putty with excellent formability properties.

REFERENCE COUNT: 20 THERE ARE 20 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L11 ANSWER 12 OF 16 HCAPLUS COPYRIGHT 2006 ACS on STN

1998:548568 HCAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 129:193756

Matrix-free osteogenic devices, implants and methods TITLE:

of use thereof

Rueger, David C.; Tucker, Marjorie M. INVENTOR(S): PATENT ASSIGNEE(S): Creative Biomolecules, Inc., USA

PCT Int. Appl., 79 pp. SOURCE:

CODEN: PIXXD2

DOCUMENT TYPE: Patent English LANGUAGE:

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 9834655 W: AU, CA, JP	A1	19980813	WO 1998-US2159	19980205 <

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RW: AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE
                                19980813
                                            CA 1998-2280931
                                                                   19980205 <--
     CA 2280931
                         AA
                                            AU 1998-62677
                                                                   19980205 <--
    AU 9862677
                          A1
                                19980826
    EP 957943
                          A1
                                19991124
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                         В1
    EP 957943
                                20030507
            AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, FI
     JP 2001511042
                          T2
                                20010807
                                            JP 1998-534870
                                                                   19980205 <--
                                                                   19980205 <--
                                            US 1998-19339
    US 6281195
                          В1
                                20010828
                                                                   19980205 <--
    AT 239514
                          Ε
                                20030515
                                            AT 1998-904920
                                            EP 2003-9751
                                                                   19980205 <--
    EP 1350525
                          A2
                                20031008
    EP 1350525
                         A3
                                20031210
            AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
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                                            US 2001-887901
                                                                   20010622 <--
     US 2002091077
                          A1
                                20020711
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     US 6426332
                          B2
    AU 779278
                          B2
                                20050113
                                            AU 2001-97272
                                                                   20011217 <--
                                                                P 19970207 <--
                                            US 1997-37327P
PRIORITY APPLN. INFO.:
                                                                P 19970529 <--
                                            US 1997-47909P
                                                                A3 19980205 <--
                                            AU 1998-62677
                                            EP 1998-904920
                                                                A3 19980205 <--
                                            US 1998-19339
                                                                A1 19980205 <--
                                                                W 19980205 <--
                                            WO 1998-US2159
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AB Provided herein are methods for inducing bone formation in a mammal sufficient to fill a defect defining a void, wherein osteogenic protein is provided alone or dispersed in a biocompatible non-rigid, amorphous carrier having no defined surfaces. The methods and devices provide injectable formulations for filling critical size defects, as well as for accelerating the rate and enhancing the quality of bone formation in non-critical size defects.

REFERÊNCE COUNT: 11 THERE ARE 11 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L11 ANSWER 13 OF 16 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1995:423960 HCAPLUS

DOCUMENT NUMBER: 122:230715

TITLE: Stimulation of osteoinduction in bone wound healing by

high-molecular hyaluronic acid

AUTHOR(S): Sasaki, T.; Watanabe, C.

CORPORATE SOURCE: School of Dentistry, Showa University, Tokyo, 142,

Japan

SOURCE: Bone (New York, NY, United States) (1995),

16(1), 9-15

CODEN: BONEDL; ISSN: 8756-3282

DOCUMENT TYPE: Journal LANGUAGE: English

AB To study the osteoinductive action of hyaluronic acid
(HA), we examined the effects of applying an elastoviscous high mol. HA
preparation on bone wound healing after bone marrow ablation. The middiaphyses
of cortical bones from rat femurs were perforated with a round bar, and
excavated marrow cavities were filled immediately with high-mol. HA. Bone
marrow ablation without HA was used to prepare controls. On post-ablation
days 1, 2, 4, 7, and 14, animals were perfusion-fixed with an aldehyde
mixture, and dissected femurs were examined by means of light, transmission-,
and scanning-electron microscopy. In controls, the wounded marrow
cavities were first filled with blood and fibrin clots (days 1 and 2),
then with granulated tissues containing macrophages, neutrophils, and
fibroblastic cells (day 4). New bone formation by differentiated
osteoblasts was observed at 1 wk post-ablation; at 2 wk, the perforated

cortical bones and marrow cavities were filled mostly with newly formed

trabecular bone. In bones to which HA had been applied, new bone formation already had been induced by day 4 on both the peri- and endosteal surfaces of the existing cortical bones. At 1 wk post-ablation, marrow cavities were completely filled with newly formed trabecular bones, in which active bone remodeling by osteoblasts and osteoclasts had occurred. Granulated tissues were replaced rapidly by normal marrow cells. These results suggest that high-mol. HA is capable of accelerating new bone formation through mesenchymal cell differentiation in bone wounds.

L11 ANSWER 14 OF 16 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1995:244750 HCAPLUS

DOCUMENT NUMBER: 122:1863

TITLE: Osteogenic protein-1, a bone morphogenic protein

member of the TGF- β superfamily, shares

chemotactic but not fibrogenic properties with

 $TGF-\beta$

AUTHOR(S): Postlethwaite, Arnold E.; Raghow, Rajendra; Stricklin,

George; Ballou, Leslie; Sampath, T. Kuber

CORPORATE SOURCE: Div. Connective Tissue Dis., Univ. Tennessee, Memphis,

TN, 38163, USA

SOURCE: Journal of Cellular Physiology (1994),

161(3), 562-70

CODEN: JCLLAX; ISSN: 0021-9541

PUBLISHER: Wiley-Liss
DOCUMENT TYPE: Journal
LANGUAGE: English

AB The authors have previously shown that recombinant human osteogenic protein-1 (rhOP-1), a bone morphogenetic protein member of the $TGF-\beta$ superfamily, can induce new bone formation when implanted with an appropriate carrier at s.c. sites in rats and can restore completely large diaphyseal segmental defects in laboratory animals. The role of OP-1 in the early events of bone induction viz, chemotaxis of phagocytic leukocytes, and fibroblastic mesenchymal cells is currently unknown. In the present study, the authors examined the effect of rhOP-1 on chemotaxis of phagocytic leukocytes (human neutrophils and monocytes) and fibroblastic mesenchymal cells (infant foreskin fibroblasts). Since OP-1 is structurally related to TGF- β 1, the authors assessed the effects of OP-1 on several other fibroblast functions (in addition to chemotaxis) known to be modulated by TGF- β 1. The results demonstrated that rhOP-1, like TGF- β 1, is a potent chemoattractant for human neutrophils, monocytes, and fibroblasts. However, in contrast to TGF- β 1, OP-1 does not to stimulate fibroblast mitogenesis, matrix synthesis [collagen and hyaluronic acid (hyaluronan)], or production of tissue inhibitor of metalloproteinase (TIMP), i.e., fibroblast functions associated with fibrogenesis. These results clearly demonstrate a dichotomy between these two members of the TGF- β superfamily with regard to fibrogenic effects on fibroblasts but a similarity in their chemotactic properties.

L11 ANSWER 15 OF 16 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1993:109744 HCAPLUS

DOCUMENT NUMBER: 118:109744

TITLE: Pharmaceutical formulations of osteogenic proteins INVENTOR(S): Ron, Eyal; Turek, Thomas J.; Isaacs, Benjamin S.;

Patel, Himakshi; Kenley, Richard A.

PATENT ASSIGNEE(S): Genetics Institute, Inc., USA

SOURCE: PCT Int. Appl., 26 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

	TENT NO.			KIN		APPLICATION NO.	DATE	
WO	9300050			A2		WO 1992-US5309	19920622 <	-
					JP, KR, NO, DK, ES, FR,	RU, US GB, GR, IT, LU, MC,	NL, SE	
AU						AU 1992-22542		-
AU	663328			B2	19951005			
EP	591392			A1	19940413	EP 1992-914339	19920622 <	-
EP	591392			В1	19960911			
	R: AT,	BE,	CH,	DE,	DK, ES, FR,	GB, GR, IT, LI, LU,	MC, NL, SE	
	06508777					JP 1993-501625	19920622 <	-
JP	3351525				20021125			
AT	142460			_	19960915	AT 1992-914339		
ES	2094359			Т3	19970116	ES 1992-914339	19920622 <	-
US	5597897			Α	19970128	US 1993-81378	19930629 <	-
NO	9304573			Α	19931213	NO 1993-4573	19931213 <	-
NO	307402			В1	20000403			
FI	109274			В1	20020628	FI 1993-5732	19931220 <	-
PRIORITY	APPLN.	INFO.	:			US 1991-718721	A 19910621 <	-
						WO 1992-US5309	A 19920622 <	-

AB Pharmaceutical formulations designed to sequester osteogenic proteins in situ for a time sufficient to allow the protein to induce cartilage and/or bone formation comprises an admixt. of an osteogenic protein, a matrix selected form the group consisting of poly(lactic acid), poly(glycolic acid), and lactic acid-glycolic acid copolymer, and an osteogenic protein-sequestering alkyl cellulose. The formulations provide malleable implants and can be used for repairing bone defects.

L11 ANSWER 16 OF 16 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1986:32407 HCAPLUS

DOCUMENT NUMBER: 104:32407

TITLE: An electron microscopic demonstration of induction of

chondrogenesis in neonatal rat muscle
outgrowth cells in monolayer cultures

AUTHOR(S): Koskinen, Kari P.; Kanwar, Yasphal S.; Sires, Bryan;

Veis, Arthur

CORPORATE SOURCE: Dent. Sch., Northwestern Univ., Chicago, IL, 60611,

USA

SOURCE: Connective Tissue Research (1985), 14(2),

141-58

CODEN: CVTRBC; ISSN: 0300-8207

DOCUMENT TYPE: Journal LANGUAGE: English

AB Second-passage fibroblast-like cells grown from explants of neonatal fetus just before normal parturition) rat muscle continue to demonstrate fibroblastlike properties for many days when cultured on plastic surfaces. Such cells can be induced to change to a chondrocytelike mode of expression by the addition of effector materials prepared from bovine cortical bone decalcified with 0.6N HCl. Other studies show that similar demineralized bone particles and exts. from them have, in vivo, osteoinductive properties. Optimum conditions for this differentiation in monolayer culture were found in the use of 2% fetal calf serum with Dulbecco's modified Eagle medium. At 10% fetal calf serum the chondrogenic changes could not be detected. Light microscopy showed a

sequence of morphol. changes, after 36 h in culture, which resembled those seen at the beginning of osteogenesis in vivo. Induced cultures showed abundant extracellular proteoglycan production. Isotope incorporation studies showed stimulation of glycosaminoglycan synthesis in response to effector materials in soluble form. Type II collagen could be detected after 3 days. Electron microscopic anal. of induced and control cultures showed unequivocal evidence for marked production of an extensive extracellular matrix in the region of effector particles. The cells themselves change shape and develop an abundant system of lysosomelike vesicles and a very active, highly engorged endoplasmic reticulum of Golgi apparatus. After 9 days in culture, evidence for the formation of a ruthenium red-stained structure on the surface of the cells in contact with inductive particles, was observed. The simple monolayer culture system described provides a direct means by which the presence of active chondrogenic fractions may be assessed, and in which the mechanism of action of the effectors can be studied.

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L13
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=> d ibib abs 113 1-6

L13 ANSWER 1 OF 6 EMBASE COPYRIGHT (c) 2006 Elsevier B.V. All rights

reserved on STN

ACCESSION NUMBER: 97365781 EMBASE

DOCUMENT NUMBER:

1997365781

TITLE:

Bone graft substitutes.

AUTHOR:

Boyan B.D.; Nasatzky E.; Keller T.A.; Schwartz Z.

CORPORATE SOURCE:

Dr. B.D. Boyan, Department of Orthopaedics, Univ. of Texas

Health Science Center, 7703 Floyd Curl Drive, San Antonio,

TX, 78284-7774, United States

SOURCE:

Current Opinion in Orthopaedics, (1997) Vol. 8,

No. 5, pp. 86-92. .

Refs: 33

ISSN: 1041-9918 CODEN: COORE

United Kingdom COUNTRY:

Journal; General Review DOCUMENT TYPE: Orthopedic Surgery FILE SEGMENT: 033

LANGUAGE: English SUMMARY LANGUAGE: English

Entered STN: 15 Jan 1998 ENTRY DATE:

Last Updated on STN: 15 Jan 1998

Bone graft substitutes span the range of processed allograft AB bone to synthetic materials and synthetic-biologic composites. past year, studies showed that there is considerable variability in the bone-inductive quality of demineralized freeze-dried bone allografts; the providers of these materials are working to find ways of ensuring quality by identifying donor characteristics that are responsible. New bone void fillers also became available. Calcium sulfate pellets were approved by the Food and Drug Administration for use in orthopedics in the United States. Among the new bone graft substitutes presented at the 1997 Orthopaedic Research Society meeting at the American Academy of Orthopedic Surgeons 1997 meeting (San Francisco, CA) were polylactic acid-polyglycolic acid scaffolds and a hyaluronic acid-basic fibroblast growth factor composite. Bone morphogenetic proteins continue to be studied both in animals and in humans. It is becoming clear that these factors may be most valuable for treatment of chronic nonunions and spine fusion. Scientists are also beginning to examine cell and gene therapies for promoting bone formation.

L13 ANSWER 2 OF 6 BIOSIS COPYRIGHT (c) 2006 The Thomson Corporation on STN

ACCESSION NUMBER: 1996:279016 BIOSIS DOCUMENT NUMBER: PREV199699001372

TITLE: Basic fibroblast growth factor for stimulation of

bone formation in osteoinductive

or conductive implants.

AUTHOR(S): Wang, Jian-Sheng

CORPORATE SOURCE: Dep. Orthopedics, Univ. Lund, Lund, Sweden SOURCE: Acta Orthopaedica Scandinavica, (1996) Vol. 67,

No. SUPPL. 269, pp. 1-33.

CODEN: AOSAAK. ISSN: 0001-6470.

DOCUMENT TYPE: Article

General Review; (Literature Review)

LANGUAGE: English

ENTRY DATE: Entered STN: 25 Jun 1996

Last Updated on STN: 25 Jun 1996

Basis Fibroblast Growth Factor (bFGF) is one of the endogenous AB factors found in bone matrix. bFGF is a mitogen for many cell types, including osteoblasts and chondrocytes. It can stimulate angiogenesis and osteoblast gene expression. The purpose of this study was to investigate whether exogenous bFGF can stimulate the formation of bone in bone grafts and in a bone graft substitute. In a model using demineralized bone matrix implants for bone induction, a dose of 15 ng bFGF per implant increased the number of chondrocytes and the amount of bone, whereas 1900 ng greatly inhibited cartilage and bone formation. These results are consistent with previous studies with this model, showing that a lower dose of bFGF increased bone calcium content and a higher dose reduced it. Thus, exogenous bFGF can stimulate proliferation during early phases of bone induction. A new device, the bone conduction chamber, was developed for the application of bFGF to bone conductive materials. This model made it possible to demonstrate a difference between the conductive properties of bone grafts and porous hydroxyapatite. bFGF increased bone ingrowth into bone graft inside the chamber and showed a biphasic dose-response curve, so that 98-200 ng per implant (0.4-10 ng/mm-3) increased bone ingrowth, but higher or lower doses had no effect. The same doses had the same effects in porous hydroxyapatite. In both bone grafts and porous hydroxyapatite, the highest dose still caused an increase in ingrowth of fibrous tissue. The effect on bone ingrowth was first detected after 6 weeks, regardless if administration of bFGF started at implantation or 2 weeks later, using an implanted minipump. Hyaluronate gel was effective as a slow-release carrier for bFGF. In conclusion, bFGF stimulates bone formation in bone implants, depending on dose and method for administration.

L13 ANSWER 3 OF 6 MEDLINE ON STN DUPLICATE 1

ACCESSION NUMBER: 95260585 MEDLINE DOCUMENT NUMBER: PubMed ID: 7742090

TITLE: Stimulation of osteoinduction in bone wound healing by

high-molecular hyaluronic acid.

AUTHOR: Sasaki T; Watanabe C

CORPORATE SOURCE: Department of Oral Anatomy, School of Dentistry, Showa

University, Tokyo, Japan.

SOURCE: Bone, (1995 Jan) Vol. 16, No. 1, pp. 9-15.

Journal code: 8504048. ISSN: 8756-3282.

PUB. COUNTRY: United States

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 199506

ENTRY DATE: Entered STN: 19950621

Last Updated on STN: 19950621 Entered Medline: 19950613

To study the osteoinductive action of hyaluronic acid AΒ (HA), we examined the effects of applying an elastoviscous high-molecular HA preparation on bone wound healing after bone marrow ablation. The middiaphyses of cortical bones from rat femurs were perforated with a round bar, and excavated marrow cavities were filled immediately with high-molecular HA. Bone marrow ablation without HA was used to prepare controls. On post-ablation days 1, 2, 4, 7, and 14, animals were perfusion-fixed with an aldehyde mixture, and dissected femurs were examined by means of light, transmission-, and scanning-electron microscopy. In controls, the wounded marrow cavities were first filled with blood and fibrin clots (days 1 and 2), then with granulated tissues containing macrophages, neutrophils, and fibroblastic cells (day 4). bone formation by differentiated osteoblasts was observed at 1 week post-ablation; at 2 weeks, the perforated cortical bones and marrow cavities were filled mostly with newly formed trabecular bone. In bones to which HA had been applied, new bone formation already had been induced by day 4 on both the peri- and endosteal surfaces of the existing cortical bones. At 1 week post-ablation, marrow cavities were completely filled with newly formed trabecular bones, in which active bone remodeling by osteoblasts and osteoclasts had occurred. Granulated tissues were replaced rapidly by normal marrow cells. These results suggest that high-molecular HA is capable of accelerating new bone formation through mesenchymal cell differentiation in bone wounds.

L13 ANSWER 4 OF 6 MEDLINE on STN DUPLICATE 2

ACCESSION NUMBER: 95051049 MEDLINE DOCUMENT NUMBER: PubMed ID: 7962137

TITLE: Osteogenic protein-1, a bone morphogenic protein member of

the TGF-beta superfamily, shares chemotactic but not

fibrogenic properties with TGF-beta.

AUTHOR: Postlethwaite A E; Raghow R; Stricklin G; Ballou L; Sampath

ΤK

CORPORATE SOURCE: Department of Medicine, University of Tennessee, Memphis

38163.

CONTRACT NUMBER: AR26034 (NIAMS)

AR39166 (NIAMS) AR39682 (NIAMS)

SOURCE: Journal of cellular physiology, (1994 Dec) Vol.

161, No. 3, pp. 562-70.

Journal code: 0050222. ISSN: 0021-9541.

PUB. COUNTRY: United States

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 199412

ENTRY DATE: Entered STN: 19950110

Last Updated on STN: 19980206 Entered Medline: 19941227

AB We have previously shown that recombinant human osteogenic protein-1 (rhOP-1), a bone morphogenetic protein member of the TGF-beta superfamily, can induce new bone formation when implanted with an appropriate carrier at subcutaneous sites in rats and can restore completely large diaphyseal segmental defects in laboratory animals. The role of OP-1 in the early events of bone induction viz, chemotaxis of phagocytic leukocytes, and fibroblastic mesenchymal cells is currently unknown. In the present study, we examined the effect of rhOP-1 on chemotaxis of phagocytic leukocytes (human neutrophils and monocytes) and

fibroblastic mesenchymal cells (infant foreskin fibroblasts). Since OP-1 is structurally related to TGF-beta 1, we assessed the effects of OP-1 on several other fibroblast functions (in addition to chemotaxis) known to be modulated by TGF-beta 1. Our results demonstrated that rhOP-1, like TGF-beta 1, is a potent chemoattractant for human neutrophils, monocytes, and fibroblasts. However, in contrast to TGF-beta 1, OP-1 does not to stimulate fibroblast mitogenesis, matrix synthesis [collagen and hyaluronic acid (hyaluronan)], or production of tissue inhibitor of metalloproteinase (TIMP), i.e., fibroblast functions associated with fibrogenesis. These results clearly demonstrate a dichotomy between these two members of the TGF-beta superfamily with a regard to fibrogenic effects on fibroblasts but a similarity in their chemotactic properties.

L13 ANSWER 5 OF 6 MEDLINE on STN DUPLICATE 3

ACCESSION NUMBER: 93386043 MEDLINE DOCUMENT NUMBER: PubMed ID: 8374499

TITLE: Bone density in old chickens' metaphyses, as affected by

local trauma and chondrocyte implantation.

AUTHOR: Robinson D; Halperin N; Nevo Z

CORPORATE SOURCE: Department of Orthopaedic Surgery, Assaf Harofeh Medical

Center, Zeriffin.

SOURCE: Bulletin (Hospital for Joint Diseases (New York, N.Y.)),

(1993 Spring) Vol. 53, No. 1, pp. 83-7. Journal code: 9215948. ISSN: 0018-5647.

PUB. COUNTRY: United States

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 199310

ENTRY DATE: Entered STN: 19931105

Last Updated on STN: 19931105 Entered Medline: 19931021

Resurfacing of joints by chondrocyte implants often leads to an increased AΒ subchondral bone density. To further evaluate this phenomenon, this study analyzed bone density and bone formation in three groups of 3-year-old chickens (90 animals, 30 per group): (1) implantation of chondrocytes embedded in hyaluronic acid (HA) into the tibial metaphysis; (2) implantation of delivery substance only; (3) sham-operated control group. Results were assessed biochemically, histologically, and histomorphometrically at 6 weeks and 6 months postimplantation. A 1.5-fold increase in the metaphyseal bone density was observed in the HA-implanted controls, as compared to sham-operated/normal joints. A further increase in bone density to twice the density of the sham-operated joints was achieved by implantation of chondrocytes. In bones implanted with cells, long-lasting (6 weeks) cartilage nodules were observed, which eventually underwent hypertrophy. The implanted chondrocytes were surrounded by a dense inflammatory infiltrate, which did not prevent the induction and formation of new bone. Based on these findings, it was concluded that chondrocyte implantation into bones results in an increase in local bone density due to a prolonged process of endochondral ossification. Further studies are necessary to evaluate the possible application of this implantation procedure in osteoporosis.

L13 ANSWER 6 OF 6 MEDLINE on STN DUPLICATE 4 ACCESSION NUMBER: 90367368 MEDLINE

DOCUMENT NUMBER: PubMed ID: 2118436

TITLE: Bone morphogenetic protein-mediated interaction of

periosteum and diaphysis. Citric acid and other factors

influencing the generation of parosteal bone.

AUTHOR: Kubler N; Urist M R

CORPORATE SOURCE: Universitatsklinik u. Polikliniken f. Zahn-, Mund- u.

Kieferkrankheiten, Wurzburg, Federal Republic of Germany.

CONTRACT NUMBER: DEO2103 (NIDCR)

SOURCE: Clinical orthopaedics and related research, (1990

Sep)_No. 258, pp. 279-94.

Journal code: 0075674. ISSN: 0009-921X.

PUB. COUNTRY: United States

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Abridged Index Medicus Journals; Priority Journals

ENTRY MONTH: 199010

ENTRY DATE: Entered STN: 19901109

Last Updated on STN: 19970203 Entered Medline: 19901005

In rabbits, after long-bone growth is complete and the cambium AB layer regresses, mesenchymal-type cells with embryonic potential (competence) for bone development persist in the adventitial layer of periosteum. These cells are not determined osteoprogenitor cells (stem cells) because bone tissue differentiation does not occur when adult periosteum is transplanted into a heterotopic site. In this respect, adventitial cells differ from bone marrow stroma cells. In a parosteal orthotopic site in the space between the adult periosteum and diaphysis, implants of bone morphogenetic protein (BMP) and associated noncollagenous proteins (BMP/NCP) induce adventitia and adjacent muscle connective-tissue-derived cells to switch from a fibrogenetic to a chondroosteoprogenetic pattern of bone development. The quantity of induced bone is proportional to the dose of BMP/NCP in the range from 10 to 50 mg; immature rabbits produced larger deposits than mature rabbits in response to BMP/NCP. Preoperative local intramuscular injections of citric, edetic, or hyaluronic acids in specified concentrations markedly enhanced subperiosteal BMP/NCP-induced bone formation. The quantity of bovine or human BMP/NCP-induced bone formation in rabbits is also increased by very low-dose immunosuppression but not by bone mineral, tricalcium phosphate ceramic, inorganic calcium salts, or various space-occupying, unspecific chemical irritants. Although composities of BMP/NCP and allogeneic rabbit tendon collagen increased the quantity of bone in a parosteal site, in a heterotopic site the composite failed to induce bone formation. In a parosteal site, the conditions permitting BMP/NCP-induced bone formation develop, and the end product of the morphogenetic response is a duplicate diaphysis. How BMP reactivates the morphogenetic process in postfetal mesenchymal-type adventitial cells persisting in adult periosteum (including adjacent muscle attachments) is not known.

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L17 ANSWER 1 OF 40 USPATFULL on STN
                      2005:326442 USPATFULL
ACCESSION NUMBER:
                      Matrix protein compositions for grafting
TITLE:
                      Lyngstadaas, St.ang.le Petter, Nesoddtangen, NORWAY
INVENTOR(S):
                      Gestrelius, STina, Lund, SWEDEN
                      Biora BioEx AB, Malmo, SWEDEN (non-U.S. corporation)
PATENT ASSIGNEE(S):
                          NUMBER KIND DATE
                      ______
                      US 6979670 B1 20051227
PATENT INFORMATION:
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APPLICATION INFO.:	US 2000-521907	20000309	(9)
	NUMBER	DATE	
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DOCUMENT TYPE: FILE SEGMENT: PRIMARY EXAMINER: LEGAL REPRESENTATIVE: NUMBER OF CLAIMS: EXEMPLARY CLAIM:	Utility GRANTED Saunders, David Kudirka & Jobse, 23	LLP	

NUMBER OF DRAWINGS: 6 Drawing Figure(s); 6 Drawing Page(s)
LINE COUNT: 1227

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB Enamel matrix, enamel matrix derivatives and/or enamel matrix proteins are used in the preparation of a pharmaceutical composition for promoting the take of a graft, e.g. in soft tissue such as skin or mucosa or mineralized tissue such as bone.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L17 ANSWER 2 OF 40 USPATFULL on STN

ACCESSION NUMBER: 2005:177362 USPATFULL

TITLE: Adipose-derived stem cells and lattices

INVENTOR(S): Katz, Adam, Charlottesville, VA, UNITED STATES

Llull, Ramon, Mallorca, SPAIN

Futrell, J. William, Pittsburgh, PA, UNITED STATES Hedrick, Marc H., Encinitas, CA, UNITED STATES Benhaim, Prosper, Encino, CA, UNITED STATES

Lorenz, Hermann Peter, Belmont, CA, UNITED STATES

NUMBER KIND

Zhu, Min, San Diego, CA, UNITED STATES Zuk, Patricia, Venice, CA, UNITED STATES Ashjian, Peter H., New York, NY, UNITED STATES

PATENT INFORMATION: US 2005153442 A1 20050714
APPLICATION INFO.: US 2004-845315 A1 20040512 (10)
RELATED APPLN. INFO.: Continuation-in-part of Ser. No. US 2003-651564, filed

on 29 Aug 2003, PENDING Continuation-in-part of Ser. No. US 2001-952522, filed on 10 Sep 2001, ABANDONED Continuation-in-part of Ser. No. US 2001-936665, filed on 10 Sep 2001, GRANTED, Pat. No. US 6777231 A 371 of International Ser. No. WO 2000-US6232, filed on 10 Mar

DATE

2000

NUMBER DATE

PRIORITY INFORMATION: US 1999-123711P 19990310 (60) <-- US 1999-162462P 19991029 (60)

DOCUMENT TYPE: Utility
FILE SEGMENT: APPLICATION

LEGAL REPRESENTATIVE: MANDEL & ADRIANO, 55 SOUTH LAKE AVENUE, SUITE 710,

PASADENA, CA, 91101, US

NUMBER OF CLAIMS: 21 EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 84 Drawing Page(s)

LINE COUNT: 9138

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The present invention provides adipose-derived stem cells (ADSCs), AB adipose-derived stem cell-enriched fractions (ADSC-EF) and adipose-derived lattices, alone and combined with the ADSCs of the invention. In one aspect, the present invention provides an ADSC substantially free of adipocytes and red blood cells and clonal populations of connective tissue stem cells. The ADSCs can be employed, alone or within biologically-compatible compositions, to generate differentiated tissues and structures, both in vivo and in vitro. Additionally, the ADSCs can be expanded and cultured to produce molecules such as hormones, and to provide conditioned culture media for supporting the growth and expansion of other cell populations. In another aspect, the present invention provides a adipose-derived lattice substantially devoid of cells, which includes extracellular matrix material from adipose tissue. The lattice can be used as a substrate to facilitate the growth and differentiation of cells, whether in vivo or in vitro, into anlagen or even mature tissues

or structures.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L17 ANSWER 3 OF 40 USPATFULL on STN

ACCESSION NUMBER: 2005:177361 USPATFULL

TITLE: Adipose-derived stem cells and lattices

INVENTOR(S): Hedrick, Marc H., Encinitas, CA, UNITED STATES Katz, Adam J., Charlottesville, VA, UNITED STATES

Llull, Ramon, Mallorca, SPAIN

Futrell, J. William, Pittsburgh, PA, UNITED STATES

Benhaim, Prosper, Encino, CA, UNITED STATES

Lorenz, Hermann Peter, Belmont, CA, UNITED STATES

Zhu, Min, Los Angeles, CA, UNITED STATES

NUMBER KIND DATE

-----PATENT INFORMATION: US 2005153441 A1 20050714 US 2003-740315 A1 20031217 (10) APPLICATION INFO.: Continuation of Ser. No. US 2001-952522, filed on 10 RELATED APPLN. INFO.:

Sep 2001, ABANDONED Continuation-in-part of Ser. No. US 2001-936665, filed on 10 Sep 2001, GRANTED, Pat. No. US 6777231 A 371 of International Ser. No. WO 2000-US6232,

filed on 10 Mar 2000

NUMBER DATE ______

US 1999-123711P 19990310 (60) PRIORITY INFORMATION: <--

US 1999-162462P 19991029 (60)

DOCUMENT TYPE: Utility APPLICATION FILE SEGMENT:

LEGAL REPRESENTATIVE: MANDEL & ADRIANO, 55 SOUTH LAKE AVENUE, SUITE 710,

PASADENA, CA, 91101, US

NUMBER OF CLAIMS: EARFFLAKY CLAIM: 1
NUMBER OF DRAWINGS: 46 Drawing Page(s)
LINE COUNT: 6606

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The present invention provides adipose-derived stem cells (ADSCs), adipose-derived stem cell-enriched fractions (ADSC-EF) and adipose-derived lattices, alone and combined with the ADSCs of the invention. In one aspect, the present invention provides an ADSC substantially free of adipocytes and red blood cells and clonal populations of connective tissue stem cells. The ADSCs can be employed, alone or within biologically-compatible compositions, to generate differentiated tissues and structures, both in vivo and in vitro. Additionally, the ADSCs can be expanded and cultured to produce molecules such as hormones, and to provide conditioned culture media for supporting the growth and expansion of other cell populations. In another aspect, the present invention provides a adipose-derived lattice substantially devoid of cells, which includes extracellular matrix material from adipose tissue. The lattice can be used as a substrate to facilitate the growth and differentiation of cells, whether in vivo or in vitro, into anlagen or even mature tissues or structures.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L17 ANSWER 4 OF 40 USPATFULL on STN

ACCESSION NUMBER: 2005:170862 USPATFULL

TITLE: Use of anti-IL-17 antibody for the treatment of

cartilage damaged by osteoarthritis

INVENTOR(S): INVENTOR(S): Filvaroff, Ellen H., San Francisco, CA, UNITED STATES PATENT ASSIGNEE(S): Genentech, Inc. (U.S. corporation)

NUMBER KIND DATE ______ PATENT INFORMATION: US 2005147609 A1 20050707 APPLICATION INFO.: US 2004-948780 A1 20040923 (10)

Continuation of Ser. No. US 2000-685823, filed on 9 Oct RELATED APPLN. INFO.:

2000, ABANDONED Continuation-in-part of Ser. No. US 1999-380142, filed on 25 Aug 1999, ABANDONED A 371 of International Ser. No. WO 1999-US10733, filed on 14 May 1999 Continuation-in-part of Ser. No. US 1999-311832,

filed on 14 May 1999, ABANDONED

NUMBER DATE -----US 2000-192103P 20000324 (60) US 1998-85579P 19980515 (60) US 1998-113621P 19981223 (60) US 1998-113621P 19981223 (60) PRIORITY INFORMATION: <--<--<--DOCUMENT TYPE: Utility APPLICATION FILE SEGMENT: LEGAL REPRESENTATIVE: GENENTECH, INC., 1 DNA WAY, SOUTH SAN FRANCISCO, CA, 94080, US 27 NUMBER OF CLAIMS: EXEMPLARY CLAIM: 1 NUMBER OF DRAWINGS: 18 Drawing Page(s) 5081 LINE COUNT: CAS INDEXING IS AVAILABLE FOR THIS PATENT. The present invention relates to methods for the treatment and repair of AB

cartilage, including cartilage damaged by injury or cartilagenous disorders, including degenerative cartilagenous disorders such as arthritis, comprising the administration of IL-17 and/or LIF antagonists (e.g., anti-IL-17 and anti-LIF antibodies). Optionally, the administration may be in combination with a cartilage agent (e.g., peptide growth factor, catabolism antagonist, osteo-, synovial, anti-inflammatory factor). Alternatively, the method provides for the treatment and repair of cartilage damaged by injury or cartilagenous disorders comprising the administration of IL-17 or LIF antagonists in combination with standard surgical techniques. Alternatively, the method provides for the treatment and repair of cartilage damaged by injury or cartilagenous disorders comprising the administration of chondrocytes previously treated with an effective amount of IL-17 and/or LIF antagonist. Alternatively, the method provides for the treatment of a mammal suffering from a cartilagenous disorder, comprising the adminstration of a therapeutically effective amount of an IL-17 and/or LIF antagonist.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L17 ANSWER 5 OF 40 USPATFULL on STN

ACCESSION NUMBER: 2005:114045 USPATFULL

TITLE: Use of a melanoma inhibiting activity factor (MIA) for

cartilage and bone repair

INVENTOR(S): Dony, Carola, Munchen, GERMANY, FEDERAL REPUBLIC OF

Proetzel, Gabriele, Schwanfeld, GERMANY, FEDERAL

REPUBLIC OF

Leser-Reiff, Ulrike, Penzberg, GERMANY, FEDERAL

REPUBLIC OF

PATENT ASSIGNEE(S): Scil Technology GmbH, Martinsried, GERMANY, FEDERAL

REPUBLIC OF (non-U.S. corporation)

NUMBER DATE

PRIORITY INFORMATION: EP 2001-99101315 19990128

DOCUMENT TYPE: Utility
FILE SEGMENT: GRANTED

DRIMARY FYAMINER: Andres

PRIMARY EXAMINER: Andres, Janet

LEGAL REPRESENTATIVE: Fulbright & Jaworski LLP

NUMBER OF CLAIMS: 13 EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 0 Drawing Figure(s); 0 Drawing Page(s)

LINE COUNT: 582

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB A melanoma inhibiting activity factor (MIA), preferably in combination with an osteoinductive protein, is a useful pharmaceutical agent for

promoting bone healing and/or cartilage repair.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L17 ANSWER 6 OF 40 USPATFULL on STN

ACCESSION NUMBER: 2005:88929 USPATFULL

TITLE: Adipose-derived stem cells and lattices

INVENTOR(S): Katz, Adam J., Charlottesville, VA, UNITED STATES

Llull, Ramon, Mallorca, SPAIN

Futrell, J. William, Pittsburgh, PA, UNITED STATES Hedrick, Marc H., Encinitas, CA, UNITED STATES Benhaim, Prosper, Encino, CA, UNITED STATES

Lorenz, Hermann Peter, Belmont, CA, UNITED STATES

Zhu, Min, San Diego, CA, UNITED STATES Zuk, Patricia, Venice, CA, UNITED STATES

Ashjian, Peter H., New York, NY, UNITED STATES

NUMBER	KIND	DATE	

PATENT INFORMATION: US 2005 APPLICATION INFO.: US 2003

US 2005076396 A1 20050407 US 2003-651564 A1 20030829 (10)

RELATED APPLN. INFO.: Continuation-in-part of Ser. No. US 2001-952522, filed on 10 Sep 2001, ABANDONED Continuation-in-part of Ser. No. US 2001-936665, filed on 10 Sep 2001, GRANTED, Pat.

No. US 6777231 Continuation-in-part of Ser. No. WO

2000-US6232, filed on 10 Mar 2000, PENDING

NUMBER DATE

PRIORITY INFORMATION: US 1999-123711P 19990310 (60) <-US 1999-162462P 19991029 (60)

DOCUMENT TYPE: Utility

FILE SEGMENT: APPLICATION
LEGAL DEPORTS TATTLE: MANDEL & ADDIANO 55 SOUTH LAKE A

LEGAL REPRESENTATIVE: MANDEL & ADRIANO, 55 SOUTH LAKE AVENUE, SUITE 710,

PASADENA, CA, 91101

NUMBER OF CLAIMS: 21 EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 64 Drawing Page(s)

LINE COUNT: 9217

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The present invention provides adipose-derived stem cells (ADSCs), adipose-derived stem cell-enriched fractions (ADSC-EF) and adipose-derived lattices, alone and combined with the ADSCs of the invention. In one aspect, the present invention provides an ADSC substantially free of adipocytes and red blood cells and clonal populations of connective tissue stem cells. The ADSCs can be employed, alone or within biologically-compatible compositions, to generate differentiated tissues and structures, both in vivo and in vitro. Additionally, the ADSCs can be expanded and cultured to produce

molecules such as hormones, and to provide conditioned culture media for supporting the growth and expansion of other cell populations. In another aspect, the present invention provides a adipose-derived lattice substantially devoid of cells, which includes extracellular matrix material from adipose tissue. The lattice can be used as a substrate to facilitate the growth and differentiation of cells, whether in vivo or in vitro, into anlagen or even mature tissues or structures.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L17 ANSWER 7 OF 40 USPATFULL on STN

ACCESSION NUMBER: 2004:248005 USPATFULL

TITLE: Methods and compositions for healing and repair of

articular cartilage

Zhang, Renwen, Rutherford, NJ, UNITED STATES INVENTOR(S):

Peluso, Diane, Marshfield, MA, UNITED STATES Morris, Elisabeth, Sherborn, MA, UNITED STATES

Genetics Institute, LLC (U.S. corporation) PATENT ASSIGNEE(S):

NUMBER KIND DATE

-----PATENT INFORMATION: US 2004192605 A1 20040930 APPLICATION INFO.: US 2004-779638 A1 20040218 (10)

Continuation of Ser. No. US 2000-493545, filed on 28 RELATED APPLN. INFO.:

Jan 2000, GRANTED, Pat. No. US 6727224

NUMBER DATE _____

US 1999-118160P 19990201 (60) PRIORITY INFORMATION: <--

DOCUMENT TYPE: Utility APPLICATION FILE SEGMENT:

LEGAL REPRESENTATIVE: FINNEGAN, HENDERSON, FARABOW, GARRETT & DUNNER, LLP,

1300 I STREET, NW, WASHINGTON, DC, 20005

NUMBER OF CLAIMS: EXEMPLARY CLAIM: 1 358 LINE COUNT:

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Methods and compositions are provided for the treatment of articular cartilage defects and disease involving the combination of tissue, such as osteochondral grafts, with active growth factor. The active growth factor is preferably a composition containing at least one bone morphogenetic protein and a suitable carrier. The method results in the regeneration of functional repair of articular cartilage tissue.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L17 ANSWER 8 OF 40 USPATFULL on STN

ACCESSION NUMBER: 2004:103728 USPATFULL

Methods and compositions for healing and repair of TITLE:

articular cartilage

Zhang, Renwen, Rutherford, NJ, United States INVENTOR(S):

Peluso, Diane, Marshfield, MA, United States Morris, Elisabeth, Sherborn, MA, United States

Genetics Institute, LLC., Cambridge, MA, United States PATENT ASSIGNEE(S):

(U.S. corporation)

NUMBER KIND DATE

Fubara 10/071,490

PATENT INFORMATION: US 6727224 B1 20040427 APPLICATION INFO.: US 2000-493545 20000128 (9)

NUMBER DATE

PRIORITY INFORMATION: US 1999-118160P 19990201 (60) <--

DOCUMENT TYPE: Utility FILE SEGMENT: GRANTED

PRIMARY EXAMINER: Low, Christopher S. F.

ASSISTANT EXAMINER: Robinson, Hope A.

LEGAL REPRESENTATIVE: Finnegan, Henderson, Farabow, Garrett & Dunner LLP.

NUMBER OF CLAIMS: 13 EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 0 Drawing Figure(s); 0 Drawing Page(s)

LINE COUNT: 390

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB Methods and compositions are provided for the treatment of articular cartilage defects and disease involving the combination of tissue, such as osteochondral grafts, with active growth factor. The active growth factor is preferably a composition containing at least one bone morphogenetic protein and a suitable carrier. The method results in the regeneration of functional repair of articular cartilage tissue.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L17 ANSWER 9 OF 40 USPATFULL on STN

ACCESSION NUMBER: 2004:2535 USPATFULL

TITLE: Osteogenic paste compositions and uses thereof

INVENTOR(S): McKay, William F., Memphis, TN, UNITED STATES

RELATED APPLN. INFO.: Continuation of Ser. No. WO 2000-US3024, filed on 4 Feb

2000, UNKNOWN

NUMBER DATE

PRIORITY INFORMATION: US 1999-118614P 19990204 (60) <--

DOCUMENT TYPE: Utility
FILE SEGMENT: APPLICATION

LEGAL REPRESENTATIVE: Kenneth A. Gandy, Woodard Emhart Naughton Moriarty &

McNett, Suite 3700, 111 Moument Circle, Indianapolis,

IN, 46204-5137

NUMBER OF CLAIMS: 29 EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 2 Drawing Page(s)

LINE COUNT: 761

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Described are osteogenic paste compositions with enhanced osteoinductive properties for use in bone repair. Compositions comprising a quickly resorbable paste carrier, a more slowly resorbed mineral matrix, and Bone Morphogenetic Protein (BMP) or other osteogenic factor are described which enable increased osteoinductive activity while retaining a reliable scaffold for the **formation** of new bone at the implant site. Methods for making and methods for therapeutic use of the compositions are also disclosed.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L17 ANSWER 10 OF 40 USPATFULL on STN

ACCESSION NUMBER: 2003:167076 USPATFULL

TITLE: Complex three-dimensional composite scaffold resistant

to delimination

Sherwood, Jill K., Edison, NJ, UNITED STATES INVENTOR(S):

Monkhouse, Donald, Radnor, PA, UNITED STATES

Gaylo, Christopher M., Princeton Junction, NJ, UNITED

STATES

PATENT ASSIGNEE(S): Therics, Inc. (U.S. corporation)

NUMBER KIND DATE ______ PATENT INFORMATION: US 2003114936 A1 20030619 US 2002-207531 A1 20020729 (10)

APPLICATION INFO.:

Continuation-in-part of Ser. No. US 1999-416346, filed RELATED APPLN. INFO.:

on 12 Oct 1999, GRANTED, Pat. No. US 6454811

NUMBER DATE _____

US 1998-103853P 19981012 (60) PRIORITY INFORMATION: <--

DOCUMENT TYPE: Utility APPLICATION FILE SEGMENT:

LEGAL REPRESENTATIVE: PATREA L. PABST, HOLLAND & KNIGHT LLP, SUITE 2000, ONE

ATLANTIC CENTER, 1201 WEST PEACHTREE STREET, N.E.,

ATLANTA, GA, 30309-3400

NUMBER OF CLAIMS: 41 EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 9 Drawing Page(s)

LINE COUNT: 2846

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The devices disclosed herein are composite implantable devices having a gradient of one or more of the following: materials, macroarchitecture, microarchitecture, or mechanical properties, which can be used to select or promote attachment of specific cell types on and in the devices prior to and/or after implantation. In preferred embodiments, the implants include complex three-dimensional structure, including curved regions and saddle-shaped areas. In various embodiments, the gradient forms a transition zone in the device from a region composed of materials or having properties best suited for one type of tissue to a region composed of materials or having properties suited for a different type of tissue. Methods to improve these devices for use in repair or replacement of cartilage and/or bone have been developed, which specifically address 1) the selection of the appropriate polymeric material for the cartilage region, 2) mechanical testing of the bone region including the effect of porosity and polymer/calcium phosphate ratio, and 3) prevention of delamination in the transition region.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L17 ANSWER 11 OF 40 USPATFULL on STN

ACCESSION NUMBER: 2003:119667 USPATFULL

TITLE: Adipose-derived stem cells and lattices Hedrick, Marc H., Encino, CA, UNITED STATES INVENTOR(S):

Katz, Adam J., Charlottesville, VA, UNITED STATES

Llull, Ramon, Mallorca, SPAIN

Futrell, J. William, Pittsburgh, PA, UNITED STATES

Benhaim, Prosper, Encino, CA, UNITED STATES

Lorenz, Hermann Peter, Belmont, CA, UNITED STATES

Zhu, Min, Los Angeles, CA, UNITED STATES

	NUMBER	KIND	DATE	
PATENT INFORMATION:	US 2003082152	A1	20030501	
APPLICATION INFO.:	US 2001-952522	A1	20010910	(9)

Continuation-in-part of Ser. No. WO 2000-US6232, filed RELATED APPLN. INFO.:

on 10 Mar 2000, UNKNOWN

NUMBER DATE _____

US 1999-123711P 19990310 (60) <--PRIORITY INFORMATION:

US 1999-162462P 19991029 (60)

Utility DOCUMENT TYPE: FILE SEGMENT: APPLICATION

MANDEL & ADRIANO, 55 SOUTH LAKE AVENUE, SUITE 710, LEGAL REPRESENTATIVE:

PASADENA, CA, 91101

NUMBER OF CLAIMS: 43 EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 43 Drawing Page(s)

LINE COUNT: 6443

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The present invention provides adipose-derived stem cells (ADSCs), adipose-derived stem cell-enriched fractions (ADSC-EF) and adipose-derivedlattices, alone and combined with the ADSCs of the invention. In one aspect, the present invention provides an ADSC substantially free of adipocytes and red blood cells and clonal populations of connective tissue stem cells. The ADSCs can be employed, alone or within biologically-compatible compositions, to generate differentiated tissues and structures, both in vivo and in vitro. Additionally, the ADSCs can be expanded and cultured to produce molecules such as hormones, and to provide conditioned culture media for supporting the growth and expansion of other cell populations. In another aspect, the present invention provides a adipose-derived lattice substantially devoid of cells, which includes extracellular matrix material from adipose tissue. The lattice can be used as a substrate to facilitate the growth and differentiation of cells, whether in vivo or in vitro, into anlagen or even mature tissues or structures.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L17 ANSWER 12 OF 40 USPATFULL on STN

ACCESSION NUMBER: 2003:67453 USPATFULL

TITLE: Resorbable scaffolds to promote cartilage regeneration Mansmann, Kevin A., 250 W. Lancaster Ave., Suite 310, INVENTOR(S):

Paoli, PA, United States 19301

NUMBER KIND DATE _____ US 6530956 B1 20030311 US 1999-393522 19990910 PATENT INFORMATION: APPLICATION INFO.: 19990910 (9)

NUMBER DATE ______

US 1998-99817P 19980910 (60) PRIORITY INFORMATION: <--

DOCUMENT TYPE: Utility GRANTED FILE SEGMENT:

PRIMARY EXAMINER: McDermott, Corrine

ASSISTANT EXAMINER: Phan, Hieu LEGAL REPRESENTATIVE: Kelly, Patrick D.

NUMBER OF CLAIMS: 24 EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 12 Drawing Figure(s); 4 Drawing Page(s)

LINE COUNT: 1690

A load-sharing resorbable scaffold is used to help transplanted AB chondrocytes or other cells generate new cartilage in a damaged joint such as a knee, hip, or shoulder. These scaffolds use two distinct matrix materials. One is a relatively stiff matrix material, designed to withstand and resist a compressive articulating load placed on the joint during the convalescent period, shortly after surgery. Due to the requirement for relatively high stiffness, this material must be denser and have less pore space than other matrices, so it will not be able to support highly rapid cell proliferation and cartilage secretion. The second material comprises a more open and porous matrix, designed to promote maximal rapid generation of new cartilage. In one preferred geometric arrangement, the stiffer matrix material is used to provide an outer rim and one or more internal runners, all of which can distribute a compressive load between them. The rim and runners create a cluster of internal cell-growing compartments, which are filled with the porous and open matrix material to encourage rapid cell reproduction and cartilage generation. These improved scaffolds can also have an articulating outer membrane with certain traits disclosed herein, bonded to and resting upon the upper edges of the runners and rim. The scaffold will support the membrane with a degree of stiffness and resiliency that allows the membrane to mimic a healthy cartilage surface. These scaffolds can be made of flexible materials, to allow them to be inserted into a damaged joint using arthroscopic methods and tools.

L17 ANSWER 13 OF 40 USPATFULL on STN

ACCESSION NUMBER: 2003:57086 USPATFULL

TITLE: Autologous immune cell therapy: cell compositions,

methods and applications to treatment of human disease

INVENTOR(S): Gruenberg, Micheal L., Poway, CA, UNITED STATES

PATENT INFORMATION: US 2003039650 A1 20030227 APPLICATION INFO.: US 2002-155404 A1 20020522 (10)

RELATED APPLN. INFO.: Continuation of Ser. No. US 1998-127138, filed on 31

Jul 1998, PENDING Division of Ser. No. US 1996-700565, filed on 25 Jul 1996, PENDING Continuation-in-part of Ser. No. WO 1996-US12170, filed on 24 Jul 1996, UNKNOWN

NUMBER DATE

PRIORITY INFORMATION: US 1995-44693P 19950726 (60) <--

DOCUMENT TYPE: Utility
FILE SEGMENT: APPLICATION

LEGAL REPRESENTATIVE: Stephanie Seidman, Heller Ehrman White & McAuliffe LLP,

7th Floor, 4350 La Jolla Village Drive, San Diego, CA,

92122

NUMBER OF CLAIMS: 25 EXEMPLARY CLAIM: 1 LINE COUNT: 2470

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB Compositions containing clinically relevant numbers of immune cells that have been isolated from a patient differentiated and/or expanded ex vivo. Methods for treating or preventing disease or otherwise altering

the immune status of the patient by reinfusing such cells into the donor are also provided. Methods for expanding and/or immune cells, including effector cells, in the absence of exogenous IL-2, and for administering the cells in the absence of co-infused IL-2 are also provided.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L17 ANSWER 14 OF 40 USPATFULL on STN

2003:44788 USPATFULL ACCESSION NUMBER: Bone morphogenic protein TITLE:

Young, Paul, Gaithersburg, MD, UNITED STATES INVENTOR(S): Ruben, Steven M., Olney, MD, UNITED STATES

Human Genome Sciences, Inc., Rockville, MD, UNITED PATENT ASSIGNEE(S):

STATES, 20850 (U.S. corporation)

NUMBER KIND DATE _____ PATENT INFORMATION:

US 2003032098 A1 20030213 US 2002-103197 A1 20020322 (10) APPLICATION INFO.:

Continuation of Ser. No. US 1999-458690, filed on 10 RELATED APPLN. INFO.:

Dec 1999, PENDING Continuation-in-part of Ser. No. WO

1999-US15783, filed on 14 Jul 1999, UNKNOWN

NUMBER DATE ______

US 1998-92922P 19980715 (60) <--PRIORITY INFORMATION:

DOCUMENT TYPE: Utility FILE SEGMENT: APPLICATION

HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE, LEGAL REPRESENTATIVE:

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS: EXEMPLARY CLAIM: 1 LINE COUNT: 8264

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The present invention relates to novel human BMP polypeptides and AΒ isolated nucleic acids containing the coding regions of the genes encoding such polypeptides. Also provided are vectors, host cells, antibodies, and recombinant methods for producing human BMP polypeptides. The invention further relates to diagnostic and

therapeutic methods useful for diagnosing and treating disorders related

to these novel human BMP polypeptides.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L17 ANSWER 15 OF 40 USPATFULL on STN

ACCESSION NUMBER: 2003:33183 USPATFULL

Device and method for regeneration and repair of TITLE:

cartilage lesions

Atkinson, Brent, Lakewood, CO, United States INVENTOR(S):

Benedict, James J., Arvada, CO, United States

Sulzer Biologics Inc., Austin, TX, United States (U.S. PATENT ASSIGNEE(S):

corporation)

NUMBER KIND DATE ______ US 6514514 B1 20030204 US 1999-250370 19990216 (9) PATENT INFORMATION:

APPLICATION INFO.:

Continuation-in-part of Ser. No. WO 1998-EP5100, filed RELATED APPLN. INFO.:

on 12 Aug 1998

NUMBER DATE

PRIORITY INFORMATION: EP 1997-810567 19970814 <--

DOCUMENT TYPE: Utility FILE SEGMENT: GRANTED

FILE SEGMENT: GRANTED
PRIMARY EXAMINER: Baker, Anne-Marie
LEGAL REPRESENTATIVE: Sheridan Ross P.C.

NUMBER OF CLAIMS: 58 EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 14 Drawing Figure(s); 8 Drawing Page(s)

LINE COUNT: 2122

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Disclosed is a cartilage repair product that induces both cell ingrowth into a bioresorbable material and cell differentiation into cartilage tissue. Such a product is useful for regenerating and/or repairing both vascular and avascular cartilage lesions, particularly articular cartilage lesions, and even more particularly mensical tissue lesions, including tears as well as segmental defects. Also disclosed is a method of regenerating and repairing cartilage lesions using such a product.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L17 ANSWER 16 OF 40 USPATFULL on STN

ACCESSION NUMBER: 2002:322566 USPATFULL

TITLE: AUTOLOGOUS IMMUNE CELL THERAPY: CELL COMPOSITIONS,

METHODS AND APPLICATIONS TO TREATMENT OF HUMAN DISEASE

INVENTOR(S): GRUENBERG, MICHEAL L., POWAY, CA, UNITED STATES

RELATED APPLN. INFO.: Division of Ser. No. US 1996-700565, filed on 25 Jul

1996, PENDING Continuation-in-part of Ser. No. WO

1996-US12170, filed on 24 Jul 1996, UNKNOWN

NUMBER DATE

PRIORITY INFORMATION: US 1995-44693P 19950726 (60) <--

DOCUMENT TYPE: Utility
FILE SEGMENT: APPLICATION

LEGAL REPRESENTATIVE: STEPHANIE SEIDMAN, HELLER EHRMAN WHITE & MCAULIFFE LLP,

7th FLOOR, 4350 LA JOLLA VILLAGE DRIVE, SAN DIEGO, CA

92122-1246, CA, 92122-1246

NUMBER OF CLAIMS: 153 EXEMPLARY CLAIM: 1 LINE COUNT: 2815

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB Compositions containing clinically relevant numbers of immune cells that have been isolated from a patient differentiated and/or expanded ex vivo. Methods for treating or preventing disease or otherwise altering the immune status of the patient by reinfusing such cells into the donor are also provided. Methods for expanding and/or immune cells, including effector cells, in the absence of exogenous IL-2, and for administering the cells in the absence of co-infused IL-2 are also provided.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L17 ANSWER 17 OF 40 USPATFULL on STN

<--

ACCESSION NUMBER:

2002:246177 USPATFULL

TITLE:

Composites for tissue regeneration and methods of

manufacture thereof

INVENTOR(S):

Sherwood, Jill K., Princeton, NJ, United States Griffith, Linda G., Cambridge, MA, United States

Brown, Scott, Princeton, NJ, United States

PATENT ASSIGNEE(S):

Massachusetts Institute of Technology, Cambridge, MA,

United States (U.S. corporation)

Therics, Inc., Princeton, NJ, United States (U.S.

corporation)

NUMBER KIND DATE ______

PATENT INFORMATION: APPLICATION INFO.:

US 6454811 B1 20020924 US 1999-416346 19991012 19991012 (9)

NUMBER DATE

PRIORITY INFORMATION:

US 1998-103853P 19981012 (60)

DOCUMENT TYPE: Utility

GRANTED FILE SEGMENT:

PRIMARY EXAMINER: McDermott, Corr ASSISTANT EXAMINER: Stewart, Alvin McDermott, Corrine LEGAL REPRESENTATIVE: Holland & Knight LLP

NUMBER OF CLAIMS: 62 EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 24 Drawing Figure(s); 4 Drawing Page(s)

LINE COUNT: 2036

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Composite devices for tissue engineering are provided having a gradient of one or more of the following: materials, macroarchitecture, microarchitecture, or mechanical properties, which can be used to select or promote attachment of specific cell types on and in the devices prior to and/or after implantation. In various embodiments, the gradient forms a transition zone in the device from a region composed of materials or having properties best suited for one type of tissue to a region composed of materials or having properties suited for a different type of tissue. The devices are made in a continuous process that imparts structural integrity as well as a unique gradient of materials in the architecture. The gradient may relate to the materials, the macroarchitecture, the microarchitecture, the mechanical properties of the device, or several of these together. The devices disclosed herein typically are made using solid free form processes, especially three-dimensional printing process (3DP.TM.). The device can be manufactured in a single continuous process such that the transition from one form of tissue regeneration scaffold and the other form of tissue regeneration scaffold have no "seams" and are not subject to differential swelling along an axis once the device is implanted into physiological fluid.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L17 ANSWER 18 OF 40 USPATFULL on STN

ACCESSION NUMBER: 2001:182096 USPATFULL

Autologous immune cell therapy: cell compositions, TITLE: methods and applications to treatment of human disease

Gruenberg, Micheal L., Poway, CA, United States INVENTOR(S):

> NUMBER KIND DATE

US 2001031253 A1 20011018 US 2001-824906 A1 20010402 (9) PATENT INFORMATION: APPLICATION INFO.:

RELATED APPLN. INFO.: Division of Ser. No. US 1996-700565, filed on 25 Jul

1996, PENDING Division of Ser. No. WO 1996-US12170,

filed on 24 Jul 1996, UNKNOWN

NUMBER DATE _____

US 1995-44693P 19950726 (60) PRIORITY INFORMATION: <--

DOCUMENT TYPE: Utility APPLICATION FILE SEGMENT:

LEGAL REPRESENTATIVE: Stephanie Seidman, Heller Ehrman White & McAuliffe LLP,

4250 Executive Square, 7th Floor, La Jolla, CA, 92037

NUMBER OF CLAIMS: EXEMPLARY CLAIM: 1
LINE COUNT: 2692

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Compositions containing clinically relevant numbers of immune cells that have been isolated from a patient differentiated and/or expanded ex vivo. Methods for treating or preventing disease or otherwise altering the immune status of the patient by reinfusing such cells into the donor are also provided. Methods for expanding and/or immune cells, including effector cells, in the absence of exogenous IL-2, and for administering

the cells in the absence of co-infused IL-2 are also provided.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L17 ANSWER 19 OF 40 USPATFULL on STN

ACCESSION NUMBER: 2000:174116 USPATFULL

Methods and compositions for programming an organic TITLE:

matrix for remodeling into a target tissue

Ashkar, Samy, Boston, MA, United States INVENTOR(S):

Atala, Anthony, Weston, MA, United States

Children's Medical Center Corporation, Boston, MA, PATENT ASSIGNEE(S):

United States (U.S. corporation)

NUMBER KIND DATE _____ US 6165487 20001226 US 1998-58048 19980409 (9) PATENT INFORMATION: APPLICATION INFO.:

Continuation-in-part of Ser. No. US 1997-937873, filed RELATED APPLN. INFO.:

on 29 Sep 1997 And a continuation of Ser. No. WO

1997-US17530, filed on 29 Sep 1997

NUMBER DATE _____

PRIORITY INFORMATION:

US 1996-27123P 19960930 (60) US 1994-27123P 19940805 (60) <--

DOCUMENT TYPE: Utility FILE SEGMENT: Granted

FILE SEGMENT: Granted PRIMARY EXAMINER: Azpuru, Carlos A.

LEGAL REPRESENTATIVE: Lahive & Cockfield, LLP, Hanley, Elizabeth A.,

Milasincic, Debra J.

LEGAL REPRESENTATIVE

Milas
NUMBER OF CLAIMS: 35
EXEMPLARY CLAIM: 1
1016

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Methods for programming a non-immunogenic matrix for remodeling into a target tissue are disclosed. Also dislosed are compositions which can promote the growth of selected tissue types in a subject.

Methods for preparing the compositions are also described. The methods and compositions are useful for treatment of tissue defects in tissues such as bone, cartilage, and muscle.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L17 ANSWER 20 OF 40 USPATFULL on STN

ACCESSION NUMBER: 1999:96274 USPATFULL

Hyaluronan based biodegradable scaffolds for tissue TITLE:

repair

Valentini, Robert F., Cranston, RI, United States INVENTOR(S):

Kim, Hyun D., Providence, RI, United States

Brown University, Providence, RI, United States (U.S. PATENT ASSIGNEE(S):

corporation)

NUMBER KIND DATE _____ US 5939323 19990817 US 1997-864709 19970528 (8) PATENT INFORMATION: APPLICATION INFO.:

> NUMBER DATE _____

PRIORITY INFORMATION: US 1996-18492P 19960528 (60) <--

DOCUMENT TYPE: Utility Granted FILE SEGMENT: PRIMARY EXAMINER: Witz, Jean C. ASSISTANT EXAMINER: Hanley, Susan

LEGAL REPRESENTATIVE: Wolf, Greenfield & Sacks, P.C.

NUMBER OF CLAIMS: 8 EXEMPLARY CLAIM: 1

2 Drawing Figure(s); 1 Drawing Page(s) NUMBER OF DRAWINGS:

848 LINE COUNT:

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

A hyaluronic acid derivitized scaffold and method of forming are disclosed. The scaffolds are useful for various

medical purposes such as tissue repair, tissue reconstruction and wound healing. In order to enhance these processes the scaffolds may be engineered to incorporate biologically active molecules such as BMP.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L17 ANSWER 21 OF 40 USPATFULL on STN

ACCESSION NUMBER: 1999:56457 USPATFULL

Cartilage induction by bone morphogenetic proteins TITLE: Hattersley, Gary, Cambridge, MA, United States INVENTOR(S):

Wolfman, Neil M., Dover, MA, United States

Morris, Elisabeth A., Southboro, MA, United States Rosen, Vicki A., Chestnut Hill, MA, United States

Genetics Institute, Inc., Cambridge, MA, United States PATENT ASSIGNEE(S):

(U.S. corporation)

NUMBER KIND DATE _____

PATENT INFORMATION: <--

US 5902785 19990511 US 1996-646193 19960507 (8) APPLICATION INFO.:

Continuation-in-part of Ser. No. US 1995-467110, filed RELATED APPLN. INFO.:

on 6 Jun 1995, now abandoned

DOCUMENT TYPE: FILE SEGMENT: Utility Granted

PRIMARY EXAMINER: Kemmerer, Elizabeth LEGAL REPRESENTATIVE: Lazar, Steven R., Gyure, Barbara A.

NUMBER OF CLAIMS: EXEMPLARY CLAIM: 1 811 LINE COUNT:

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Compositions of proteins with cartilaginous tissue inducing and maintenance activity are disclosed. The compositions are useful in the treatment of osteoarthritis, cartilage defects and in related tissue

repair.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L17 ANSWER 22 OF 40 USPATFULL on STN

1999:53151 USPATFULL ACCESSION NUMBER:

Bone-derived implant for load-supporting applications TITLE:

Boyce, Todd M., Aberdeen, NJ, United States INVENTOR(S): Manrique, Albert, Manalapan, NJ, United States Scarborough, Nelson L., Ocean, NJ, United States Russell, James L., Little Silver, NJ, United States Osteotech, Inc., Eatontown, NJ, United States (U.S. PATENT ASSIGNEE(S):

corporation)

NUMBER KIND DATE -----

US 5899939 19990504 US 1998-9997 19980121 (9) <--PATENT INFORMATION:

APPLICATION INFO.:

DOCUMENT TYPE: Utility FILE SEGMENT: Granted

PRIMARY EXAMINER: Granted Prebilic, Paul B. LEGAL REPRESENTATIVE: Dilworth & Barrese

NUMBER OF CLAIMS: 33 1,28 EXEMPLARY CLAIM:

NUMBER OF DRAWINGS: 6 Drawing Figure(s); 3 Drawing Page(s)

LINE COUNT: 484

A bone-derived implant is provided which is made up of one or more layers of fully mineralized or partially demineralized certical bone and, optionally, one or more layers of some other material. The layers constituting the implant are assembled into a unitary structure to provide an implant exhibiting good overall load-supporting properties.

L17 ANSWER 23 OF 40 USPATFULL on STN

ACCESSION NUMBER: 1999:36949 USPATFULL Engineering oral tissues TITLE:

Mooney, David J., Ann Arbor, MI, United States INVENTOR(S):

Rutherford, Robert B., Ann Arbor, MI, United States

The Regents of the University of Michigan, Ann Arbor, PATENT ASSIGNEE(S):

MI, United States (U.S. corporation)

NUMBER KIND DATE _____

US 5885829 19990323 <--PATENT INFORMATION: US 1997-864494 19970528 (8) APPLICATION INFO.:

> NUMBER DATE _____

US 1996-18450P 19960528 (60) <--PRIORITY INFORMATION:

DOCUMENT TYPE: Utility FILE SEGMENT: Granted

PRIMARY EXAMINER:

Degen, Nancy

LEGAL REPRESENTATIVE:

Arnold, White & Durkee

NUMBER OF CLAIMS:

109 1

EXEMPLARY CLAIM: NUMBER OF DRAWINGS:

17 Drawing Figure(s); 11 Drawing Page(s)

LINE COUNT:

8001

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB

Disclosed are methods for regenerating dental and oral tissues from viable cells using ex vivo culture on a structural matrix. The regenerated oral tissues and tissue-matrix preparations thus provided have both clinical applications in dentistry and oral medicine and are also useful in in vitro toxicity and biocompatibility testing.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L17 ANSWER 24 OF 40 USPATFULL on STN

ACCESSION NUMBER:

1999:12772 USPATFULL

TITLE: INVENTOR(S):

Nucleic acids encoding osteogenic proteins Oppermann, Hermann, Medway, MA, United States Ozkaynak, Engin, Milford, MA, United States

Kuberasampath, Thangavel, Medway, MA, United States Rueger, David C., Hopkinton, MA, United States

Pang, Roy H. L., Medway, MA, United States

PATENT ASSIGNEE(S):

Stryker Corporation, Kalamazoo, MI, United States (U.S.

corporation)

		NUMBER	KIND	DATE	
,	TNEODMATION.	HC 5863758		10000126	

PATENT INFORMATION: APPLICATION INFO.: RELATED APPLN. INFO.:

<--US 1994-449700 US 5863758 19990126 19940523 (8)

Division of Ser. No. US 1993-147023, filed on 1 Nov 1993, now patented, Pat. No. US 5468845 which is a division of Ser. No. US 1992-841646, filed on 21 Feb 1992, now patented, Pat. No. US 5266683 which is a continuation-in-part of Ser. No. US 1992-827052, filed on 28 Jan 1992, now patented, Pat. No. US 5250302 Ser. No. Ser. No. US 1990-579865, filed on 7 Sep 1990, now patented, Pat. No. US 5108753 Ser. No. Ser. No. US 1990-621849, filed on 4 Dec 1990, now abandoned Ser. No. Ser. No. US 1990-621988, filed on 4 Dec 1990, now abandoned Ser. No. Ser. No. US 1991-810560, filed on 20 Dec 1991, now abandoned Ser. No. Ser. No. US 1990-569920, filed on 20 Aug 1990, now abandoned Ser. No. Ser. No. US 1990-600024, filed on 18 Oct 1990, now abandoned Ser. No. Ser. No. US 1990-599543, filed on 18 Oct 1990, now abandoned Ser. No. Ser. No. US 1990-616374, filed on 21 Nov 1990, now patented, Pat. No. US 5162114 And Ser. No. US 1990-483913, filed on 22 Feb 1990, now patented, Pat. No. US 5171574 , said Ser. No. US 827052 which is a division of Ser. No. US 1988-179406, filed on 8 Apr 1988, now patented, Pat. No. US 4968590 , said Ser. No. US 579865 which is a division of Ser. No. US 179406, said Ser. No. US 621849 which is a division of Ser. No. US 1988-232630, filed on 15 Aug 1988, now abandoned which is a continuation-in-part of Ser. No. US 179406, said Ser. No. US 621988 which is a division of Ser. No. US 1989-315342, filed on 23 Feb 1989, now patented, Pat. No. US 5011691 which is a continuation-in-part of Ser. No. US 232630 , said Ser. No. US 810560 which is a

continuation of Ser. No. US 1991-660162, filed on 22 Feb 1991, now abandoned which is a continuation of Ser. No. US 1989-422699, filed on 17 Oct 1989, now abandoned which is a continuation-in-part of Ser. No. US 315342, said Ser. No. US 569920 which is a continuation-in-part of Ser. No. US 422699 And Ser. No. US 483913 which is a continuation-in-part of Ser. No. US 1989-422613, filed on 17 Oct 1989, now patented, Pat. No. US 4975526 which is a continuation-in-part of Ser. No. US 315342 , said Ser. No. US 600024 which is a continuation-in-part of Ser. No. US 569920 , said Ser. No. US 599543 which is a continuation-in-part of Ser. No. US 569920 , said Ser. No. US 616374 which is a division of Ser. No. US 422613

DOCUMENT TYPE: FILE SEGMENT:

Utility Granted

PRIMARY EXAMINER:

Loring, Susan A.

LEGAL REPRESENTATIVE:

Testa, Hurwitz & Thibeault, LLP

NUMBER OF CLAIMS:

49

EXEMPLARY CLAIM: NUMBER OF DRAWINGS:

80 Drawing Figure(s); 49 Drawing Page(s)

5104 LINE COUNT:

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Disclosed are (1) osteogenic devices comprising a matrix containing AB substantially pure natural-sourced mammalian osteogenic protein; (2) DNA and amino acid sequences for novel polypeptide chains useful as subunits of dimeric osteogenic proteins; (3) vectors carrying sequences encoding these novel polypeptide chains and host cells transfected with these vectors; (4) methods of producing these polypeptide chains using recombinant DNA technology; (5) antibodies specific for these novel polypeptide chains; (6) osteogenic devices comprising these recombinantly produced proteins in association with an appropriate carrier matrix; and (7) methods of using the osteogenic devices to mimic the natural course of endochondral bone formation in mammals.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L17 ANSWER 25 OF 40 USPATFULL on STN

ACCESSION NUMBER:

1998:162472 USPATFULL

TITLE:

Compositions and therapeutic methods using morphogenic

proteins and stimulatory factors

INVENTOR(S):

Lee, John C., San Antonio, TX, United States

Yeh, Lee-Chuan C., San Antonio, TX, United States

PATENT ASSIGNEE(S):

Stryker Corporation, Kalamazoo, MI, United States (U.S.

corporation)

NUMBER KIND DATE ______

PATENT INFORMATION:

US 5854207 19981229

APPLICATION INFO.:

19980223

RELATED APPLN. INFO.:

US 1998-27873 Division of Ser. No. US 1995-570752, filed on 12 Dec

1995 Utility

DOCUMENT TYPE: FILE SEGMENT:

Granted

PRIMARY EXAMINER:

Nutter, Nathan M.

LEGAL REPRESENTATIVE:

Fish & Neave, Haley, Jr., James F., Ruskin, Barbara A.

NUMBER OF CLAIMS: EXEMPLARY CLAIM:

28

NUMBER OF DRAWINGS:

12 Drawing Figure(s); 12 Drawing Page(s)

LINE COUNT:

3072

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

<--

The present invention provides pharmaceutical compositions comprising a AB morphogenic protein stimulatory factor (MPSF) for improving the tissue inductive activity of morphogenic proteins, particularly those belonging to the BMP protein family. Methods for improving the tissue inductive activity of a morphogenic protein in a mammal using those compositions are provided. This invention also provides implantable morphogenic devices comprising a morphogenic protein and a MPSF disposed within a carrier, that are capable of inducing tissue formation in allogeneic and xenogeneic implants. Methods for inducing local tissue formation from a progenitor cell in a mammal using those devices are also provided. A method for accelerating allograft repair in a mammal using morphogenic devices is provided. This invention also provides a prosthetic device comprising a prosthesis coated with a morphogenic protein and a MPSF, and a method for promoting in vivo integration of an implantable prosthetic device to enhance the bond strength between the prosthesis and the existing target tissue at the joining site. Methods of treating tissue degenerative conditions in a mammal using the pharmaceutical compositions are also provided.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L17 ANSWER 26 OF 40 USPATFULL on STN

1998:154240 USPATFULL ACCESSION NUMBER:

TITLE:

Compositions comprising bone morphogenic proteins and

truncated parathyroid hormone related peptide and

methods of inducing cartilage by administration of same Hattersley, Gary, 10 Rogers St., #303, Cambridge, MA,

INVENTOR(S): United States 02142

Rosen, Vicki A., 2 Cedar Rd., Chestnut Hill, MA, United

States 02167

NUMBER KIND DATE ______

US 5846931 19981208 US 1997-926942 19970910 (8) <--PATENT INFORMATION:

APPLICATION INFO.:

Continuation of Ser. No. US 1996-622101, filed on 26 RELATED APPLN. INFO.:

Mar 1996, now patented, Pat. No. US 5700774

DOCUMENT TYPE: Utility FILE SEGMENT: Granted

PRIMARY EXAMINER: Kemmerer, Elizabeth LEGAL REPRESENTATIVE: Lazar, Steven R.

NUMBER OF CLAIMS: EXEMPLARY CLAIM: LINE COUNT: 637

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Compositions of proteins with chondrocyte and cartilaginous tissue inducing activity, as well as method of using those compositions, are disclosed. The compositions comprise one or more proteins of the transforming growth factor- β (TGF- β)

superfamily of proteins, particularly bone morphogenetic proteins (BMPs), in combination with parathyroid hormone related polypeptide (PTHrP) or an equivalent PTH-like polypeptide. The compositions and methods are useful in the treatment of osteoarthritis, cartilage defects and in related tissue repair.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L17 ANSWER 27 OF 40 USPATFULL on STN

ACCESSION NUMBER: 1998:149524 USPATFULL

TITLE: Method for repairing cartilage

Naughton, Gail K., Del Mar, CA, United States INVENTOR(S):

Willoughby, Jane, Solana Beach, CA, United States

Advanced Tissue Sciences, Inc., La Jolla, CA, United PATENT ASSIGNEE(S):

States (U.S. corporation)

NUMBER KIND DATE ______

US 5842477 19981201 US 1996-604284 19960221 (8) <--PATENT INFORMATION:

APPLICATION INFO .:

DOCUMENT TYPE: Utility FILE SEGMENT: Granted

PRIMARY EXAMINER: Prebilic, Paul B. Pennie & Edmonds LLP LEGAL REPRESENTATIVE:

NUMBER OF CLAIMS: 48 EXEMPLARY CLAIM: 1467 LINE COUNT:

The present invention relates to methods of making and/or repairing AB cartilage in vivo comprising implanting into a patient, at a site of cartilage damage or loss, a biocompatible, non-living three-dimensional scaffold or framework structure in combination with periosteal/perichondrial tissue that can be used to hold the scaffold in place and provides a source of chondrocyte progenitor cells,

chondrocytes and other stromal cells for attachment to the scaffold in vivo. In addition, a preparation of cells that can include chondrocytes, chondrocyte progenitor cells or other stromal cells is administered, either before, during or after implantation of the scaffold and/or the periosteal perichondrial tissue; the cells are administered directly into the site of the implant in vivo and promote the induction of factors that enhance chondrogenesis and the migration of chondrocytes, progenitor cells and other stromal cells from the adjacent in vivo environment into the scaffold for the production of new cartilage at the

site of implantation.

L17 ANSWER 28 OF 40 USPATFULL on STN

1998:144150 USPATFULL ACCESSION NUMBER:

Semi-interpenetrating polymer networks TITLE:

Shastri, Venkatram R., Allston, MA, United States INVENTOR(S):

> Langer, Robert S., Newton, MA, United States Tarcha, Peter J., Lake Villa, IL, United States

Massachusetts Institute of Technology, Cambridge, MA, PATENT ASSIGNEE(S):

United States (U.S. corporation)

NUMBER KIND DATE _____

US 5837752 19981117 US 1997-895762 19970717 (8) <--PATENT INFORMATION: APPLICATION INFO.:

DOCUMENT TYPE: Utility FILE SEGMENT: Granted

PRIMARY EXAMINER: Buttner, David

Arnall Golden & Gregory, LLP LEGAL REPRESENTATIVE:

NUMBER OF CLAIMS: EXEMPLARY CLAIM: 1 927 LINE COUNT:

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Compositions for bone repair have been developed based on linear hydrophobic degradable polymers and monomers or macromers, at least one of which includes an anhydride linkage. The monomers and/or macromers crosslink each other but not to the linear polymer to form semi-interpenetrating networks. The compositions can include various

excipients, therapeutic and/or diagnostic agents. The compositions can be polymerized in the presence of dissolvable particles such as inorganic salts and proteinaceous materials to provide a porous polymer network. The compositions can be injected into a patient and polymerized in situ or can be polymerized ex vivo and implanted. When polymerized ex vivo, the composition can be shaped into various articles, such as pins, screws, and hollow tubes, which can be used to repair broken bones.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

USPATFULL on STN L17 ANSWER 29 OF 40

1998:12129 USPATFULL ACCESSION NUMBER:

Method of selectively extracting osteogenic protein TITLE:

Oppermann, Hermann, Medway, MA, United States INVENTOR(S):

Ozkaynak, Engin, Milford, MA, United States Kuberasampath, Thangavel, Medway, MA, United States

Rueger, David C., Hopkinton, MA, United States

Pang, Roy H. L., Medway, MA, United States

Stryker Corporation, Natick, MA, United States (U.S. PATENT ASSIGNEE(S):

NUMBER KIND DATE

corporation)

PATENT INFORMATION: APPLICATION INFO.: RELATED APPLN. INFO.:

______ US 5714589 19980203 US 1995-447570 19950523 (8) <--Division of Ser. No. US 1993-147023, filed on 1 Nov 1993, now patented, Pat. No. US 5468845 which is a division of Ser. No. US 1992-841646, filed on 21 Feb 1992, now patented, Pat. No. US 5266683 which is a continuation-in-part of Ser. No. US 1992-827052, filed on 28 Jan 1992, now patented, Pat. No. US 5250302 Ser. No. Ser. No. US 1990-579865, filed on 7 Sep 1990, now

patented, Pat. No. US 5108753 Ser. No. Ser. No. US 1990-621849, filed on 4 Dec 1990, now abandoned Ser. No. Ser. No. US 1990-621988, filed on 4 Dec 1990, now abandoned Ser. No. Ser. No. US 1991-810560, filed on 20 Dec 1991, now abandoned Ser. No. Ser. No. US 1990-569920, filed on 20 Aug 1990, now abandoned Ser. No. Ser. No. US 1990-600024, filed on 18 Oct 1990, now

abandoned Ser. No. Ser. No. US 1990-599543, filed on 18 Oct 1990, now abandoned Ser. No. Ser. No. US 1990-616374, filed on 21 Nov 1990, now patented, Pat. No. US 5162114 And Ser. No. US 1990-483913, filed on 22

Feb 1990, now patented, Pat. No. US 5171574, said Ser. -827052 which is a division of Ser. No. US 1988-179406, filed on 8 Apr 1988, now patented, Pat. No. US 4968590 , said Ser. No. US -579865 which is a division of Ser. No. US -179406, said Ser. No. US

-621849 which is a division of Ser. No. US 1988-232630, filed on 15 Aug 1988, now abandoned which is a -179406 , said continuation-in-part of Ser. No. US

Ser. No. US -621988 which is a division of Ser. No. US 1989-315342, filed on 23 Feb 1989, now patented, Pat. No. US 5011691 which is a continuation-in-part of -232630 , said Ser. No. US Ser. No. US -810560

which is a continuation of Ser. No. US 1991-660162, filed on 22 Feb 1991, now abandoned which is a continuation of Ser. No. US 1989-422699, filed on 17 Oct 1989, now abandoned which is a continuation-in-part

-315342 , said Ser. No. US of Ser. No. US

which is a continuation-in-part of Ser. No. US 1989-422699, filed on 17 Oct 1989 And Ser. No. US -483913 which is a continuation-in-part of Ser. No. US 1989-422613, filed on 17 Oct 1989, now patented, Pat. No. US 4975526 which is a continuation-in-part of Ser. -315342 , said Ser. No. US -600024 which is No. US a continuation-in-part of Ser. No. US -569920 , said Ser. No. US -599543 which is a continuation-in-part of Ser. No. US -569920 , said Ser. No. US

which is a division of Ser. No. US -422613

DOCUMENT TYPE: Utility FILE SEGMENT: Granted

PRIMARY EXAMINER: Loring, Susan A.

LEGAL REPRESENTATIVE: Testa, Hurwitz & Thibeault, LLP

NUMBER OF CLAIMS: 16 EXEMPLARY CLAIM:

NUMBER OF DRAWINGS: 80 Drawing Figure(s); 49 Drawing Page(s)

LINE COUNT: 4132

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Disclosed are (1) osteogenic devices comprising a matrix containing substantially pure natural-sourced mammalian osteogenic protein; (2) DNA and amino acid sequences for novel polypeptide chains useful as subunits of dimeric osteogenic proteins; (3) vectors carrying sequences encoding these novel polypeptide chains and host cells transfected with these vectors; (4) methods of producing these polypeptide chains using recombinant DNA technology; (5) antibodies specific for these novel polypeptide chains; (6) osteogenic devices comprising these recombinantly produced proteins in association with an appropriate carrier matrix; and (7) methods of using the osteogenic devices to mimic the natural course of endochondral bone formation in mammals.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L17 ANSWER 30 OF 40 USPATFULL on STN

ACCESSION NUMBER: 97:120591 USPATFULL

Compositions comprising bone morphogenic proteins and TITLE:

truncated parathyroid hormone related peptide, and

methods of inducing cartilage by administration of same INVENTOR(S): Hattersley, Gary, Cambridge, MA, United States

Rosen, Vicki A., Chestnut Hill, MA, United States

Genetics Institute, Inc., Cambridge, MA, United States PATENT ASSIGNEE(S):

(U.S. corporation)

NUMBER KIND DATE ______

PATENT INFORMATION: <--

US 5700774 19971223 US 1996-622101 19960326 (8) APPLICATION INFO.:

DOCUMENT TYPE: Utility FILE SEGMENT: Granted

Fitzgerald, David L. PRIMARY EXAMINER: Kemmerer, Elizabeth C. ASSISTANT EXAMINER: Meinert, M. C., Lazar, S. LEGAL REPRESENTATIVE:

NUMBER OF CLAIMS: 17 EXEMPLARY CLAIM: 1 668 LINE COUNT:

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Compositions of proteins with chondrocyte and cartilaginous tissue AB inducing activity, as well as method of using those compositions, are disclosed. The compositions comprise one or more proteins of the transforming growth factor- β (TGF- β)

superfamily of proteins, particularly bone morphogenetic proteins (BMPs), in combination with parathyroid hormone related polypeptide (PTHrP) or an equivalent PTH-like polypeptide. The compositions and methods are useful in the treatment of osteoarthritis, cartilage defects and in related tissue repair.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L17 ANSWER 31 OF 40 USPATFULL on STN

ACCESSION NUMBER: 97:120112 USPATFULL

TITLE: Tissue-engineered bone repair using cultured periosteal

cells

Breitbart, Arnold S., Great Neck, NY, United States INVENTOR(S):

Grande, Daniel A., Sea Cliff, NY, United States

North Shore University Hospital Research Corporation, PATENT ASSIGNEE(S):

Manhasset, NY, United States (U.S. corporation)

NUMBER KIND DATE -----

US 5700289 19971223 US 1995-545988 19951020 (8) PATENT INFORMATION: <--

APPLICATION INFO.:

Utility DOCUMENT TYPE: FILE SEGMENT: Granted

PRIMARY EXAMINER: Brittingham, Debra S.

LEGAL REPRESENTATIVE: Arnall Golden & Gregory, LLP

NUMBER OF CLAIMS: 15 EXEMPLARY CLAIM: 1

6 Drawing Figure(s); 3 Drawing Page(s) NUMBER OF DRAWINGS:

LINE COUNT: 856

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Periosteal cells have been grown in cell culture and have been shown to have an osteoblastic phenotype, with production of osteocalcin and glycosaminoglycan. When seeded into polymeric implants, repair of critical size cranial defects was demonstrated and was confirmed by histology, biochemical assays, and radiodensitometry.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L17 ANSWER 32 OF 40 USPATFULL on STN

ACCESSION NUMBER: . 97:90970 USPATFULL

Terminally sterilized osteogenic devices and TITLE:

preparation thereof

INVENTOR(S): Tucker, Marjorie M., Holliston, MA, United States

Rueger, David C., Southborough, MA, United States Sampath, Kuber T., Holliston, MA, United States

Stryker Corporation, Kalamazoo, MI, United States (U.S. PATENT ASSIGNEE(S):

corporation)

NUMBER KIND DATE _______

US 5674292 US 1995-478452 19971007 PATENT INFORMATION: <--APPLICATION INFO.: 19950607 (8)

Utility DOCUMENT TYPE: FILE SEGMENT: Granted

Kulkosky, Peter F. PRIMARY EXAMINER:

Testa, Hurwitz & Thibeault, LLP LEGAL REPRESENTATIVE:

NUMBER OF CLAIMS: 23 EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 1 Drawing Figure(s); 1 Drawing Page(s)

LINE COUNT:

Disclosed are terminally sterilized osteogenic devices for implantation AB into a mammal. The devices contain a combination of a biologically active osteogenic protein and an insoluble carrier which after being combined are sterilized by exposure to ionizing radiation, for example, by exposure to gamma rays or an electron beam. The terminally sterilized devices of the invention are characterized in that they induce bone formation following implantation into a mammal. Also disclosed is a method for inducing bone formation in a mammal by implanting a terminally sterilized device of the invention into a preselected locus in a mammal. Also disclosed is a method for preparing the terminally sterilized device of the invention.

L17 ANSWER 33 OF 40 USPATFULL on STN

ACCESSION NUMBER: 97:35731 USPATFULL

Prosthetic articular cartilage TITLE:

Stone, Kevin R., Mill Valley, CA, United States INVENTOR(S):

Li, Shu-Tung, Oakland, NJ, United States

ReGen Biologics, Inc., Redwood City, CA, United States PATENT ASSIGNEE(S):

(U.S. corporation)

NUMBER KIND DATE ______

US 5624463 19970429 US 1994-232743 19940425 (8) PATENT INFORMATION:

APPLICATION INFO.:

RELATED APPLN. INFO.: Continuation of Ser. No. US 1991-809003, filed on 17

Dec 1991 which is a continuation-in-part of Ser. No. US

1990-520027, filed on 7 May 1990 which is a

continuation-in-part of Ser. No. US 1989-317851, filed on 2 Mar 1989 which is a continuation-in-part of Ser.

No. US 1987-75352, filed on 20 Jul 1987

DOCUMENT TYPE: Utility Granted FILE SEGMENT:

Isabella, David PRIMARY EXAMINER:

LEGAL REPRESENTATIVE: Fish & Richardson P.C.

22 NUMBER OF CLAIMS: EXEMPLARY CLAIM: 1

12 Drawing Figure(s); 5 Drawing Page(s) NUMBER OF DRAWINGS:

LINE COUNT: 962

A prosthetic, resorbable articular cartilage and methods of its AR fabrication and insertion are disclosed. The prosthetic articular cartilage, when implanted in a humanoid joint, acts as a resorbable scaffold for ingrowth of native articular chondrocytes and supports natural articulating joint forces. The prosthetic articular cartilage is a dry, porous, volume matrix of biocompatible and bioresorbable fibers. These fibers include a natural polymer or analogs thereof, at least a portion of which may be crosslinked. The matrix is adapted to have an in vivo outer surface contour substantially the same as that of natural articular cartilage in an articulating joint, and has a pore size in the approximate range of about 100 microns to about 400 microns.

L17 ANSWER 34 OF 40 USPATFULL on STN

95:103607 USPATFULL ACCESSION NUMBER:

Antibodies to osteogenic proteins TITLE:

Oppermann, Hermann, Medway, MA, United States INVENTOR(S):

Ozkaynak, Engin, Milford, MA, United States

Kuberasampath, Thangavel, Medway, MA, United States

<--

PATENT ASSIGNEE(S):

Rueger, David C., Hopkinton, MA, United States Stryker Corporation, Kalamazoo, MI, United States (U.S. corporation)

NUMBER	KIND	DATE

PATENT INFORMATION: APPLICATION INFO.: RELATED APPLN. INFO.: US 5468845 19951121 US 1993-147023 19931101 (8)

Division of Ser. No. US 1992-841646, filed on 21 Feb 1992, now patented, Pat. No. US 5266683 which is a continuation-in-part of Ser. No. US 1992-827052, filed on 28 Jan 1992, now patented, Pat. No. US 5250302 Ser. No. Ser. No. US 1990-579865, filed on 7 Sep 1990, now patented, Pat. No. US 5108153 Ser. No. Ser. No. US 1990-621849, filed on 4 Dec 1990, now abandoned Ser. No. Ser. No. US 1990-621988, filed on 4 Dec 1990, now abandoned Ser. No. Ser. No. US 1991-810560, filed on 20 Dec 1991, now abandoned Ser. No. Ser. No. US 1990-569920, filed on 20 Aug 1990, now abandoned Ser. No. Ser. No. US 1990-600024, filed on 18 Oct 1990, now abandoned Ser. No. Ser. No. US 1990-599543, filed on 18 Oct 1990, now abandoned Ser. No. Ser. No. US 1990-616374, filed on 21 Nov 1990, now patented, Pat. No. US 5162114 And Ser. No. US 1990-483913, filed on 22 Feb 1990, now patented, Pat. No. US 5171594 , said Ser. -827052 which is a division of Ser. No. US No. US 1988-179406, filed on 8 Apr 1988, now patented, Pat. No. US 4968590 , said Ser. No. US 1990-579865, filed on 7 Sep 1990, now patented, Pat. No. US 5108153 which is a division of Ser. No. US -179406 , said Ser. No. US -621849 which is a division of Ser. No. US 1988-232630, filed on 15 Aug 1988, now abandoned which is a continuation-in-part of Ser. No. US -179406 , said Ser. No. US -621988 which is a division of Ser. No. US 1989-315342, filed on 23 Feb 1989, now patented, Pat. No. US 5011691 which is a continuation-in-part of -232630 , said Ser. No. US Ser. No. US which is a continuation of Ser. No. US 1991-660162, filed on 22 Feb 1991, now abandoned which is a continuation of Ser. No. US 1989-422699, filed on 17 Oct 1989, now abandoned which is a continuation-in-part -315342 , said Ser. No. US -569920 of Ser. No. US which is a continuation-in-part of Ser. No. US -422699 And Ser. No. US -483913 which is a continuation-in-part of Ser. No. US 1989-422613, filed on 17 Oct 1989, now patented, Pat. No. US 4975526 which -315342 , is a continuation-in-part of Ser. No. US said Ser. No. US 1990-600024, filed on 18 Oct 1990, now abandoned which is a continuation-in-part of Ser. No. -569920 , said Ser. No. US -599543 , said Ser. US -599543 No. US

DOCUMENT TYPE: FILE SEGMENT: PRIMARY EXAMINE

LINE COUNT:

PRIMARY EXAMINER:
ASSISTANT EXAMINER:
LEGAL REPRESENTATIVE:

NUMBER OF CLAIMS: EXEMPLARY CLAIM: NUMBER OF DRAWINGS:

Loring, Susan A. Testa, Hurwitz & Thibeault

Utility

Granted

Lacey, David L.

5 1

80 Drawing Figure(s); 49 Drawing Page(s) 4082

Searched by Mary Jane Ruhl Ext. 22524

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Disclosed are (1) osteogenic devices comprising a matrix containing AB substantially pure natural-sourced mammalian osteogenic protein; (2) DNA and amino acid sequences for novel polypeptide chains useful as subunits of dimeric osteogenic proteins; (3) vectors carrying sequences encoding these novel polypeptide chains and host cells transfected with these vectors; (4) methods of producing these polypeptide chains using recombinant DNA technology; (5) antibodies specific for these novel polypeptide chains; (6) osteogenic devices comprising these recombinantly produced proteins in association with an appropriate carrier matrix; and (7) methods of using the osteogenic devices to mimic the natural course of endochondral bone formation in mammals.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L17 ANSWER 35 OF 40 USPATFULL on STN

ACCESSION NUMBER: 94:88501 USPATFULL TITLE: Osteogenic devices

Oppermann, Hermann, Medway, MA, United States INVENTOR(S):

Ozkaynak, Engin, Milford, MA, United States Kuberasampath, Thangavel, Medway, MA, United States

Rueger, David C., Hopkinton, MA, United States

Pang, Roy H. L., Medway, MA, United States PATENT ASSIGNEE(S):

Stryker Corporation, Kalamazoo, MI, United States (U.S.

corporation)

NUMBER KIND DATE _______ US 5354557 19941011 <--US 1992-993387 19921218 (7)

PATENT INFORMATION: APPLICATION INFO.: DISCLAIMER DATE: RELATED APPLN. INFO.:

20080430 Division of Ser. No. US 1992-841646, filed on 21 Feb 1992, now patented, Pat. No. US 5266683 which is a continuation-in-part of Ser. No. US 1992-827052, filed on 28 Jan 1992, now patented, Pat. No. US 5250302 which is a division of Ser. No. US 1988-179406, filed on 8 Apr 1988, now patented, Pat. No. US 4968590 And a continuation-in-part of Ser. No. US 1990-579865, filed on 7 Sep 1990, now patented, Pat. No. US 5108753 which is a division of Ser. No. US 1988-179406, filed on 8 Apr 1988, now patented, Pat. No. US 4968590 And a continuation-in-part of Ser. No. US 1990-621849, filed on 4 Dec 1990, now abandoned which is a division of Ser. No. US 1988-232630, filed on 15 Aug 1988 which is a continuation-in-part of Ser. No. US 1988-179406, filed on 8 Apr 1988, now patented, Pat. No. US 4968590 And a continuation-in-part of Ser. No. US 1990-621988, filed on 4 Dec 1990 which is a division of Ser. No. US 1989-315342, filed on 23 Feb 1989, now patented, Pat. No. US 5011691 which is a continuation-in-part of Ser. No. US 1988-232630, filed on 15 Aug 1988 And a continuation-in-part of Ser. No. US 1991-810560, filed on 20 Dec 1991, now abandoned which is a continuation of Ser. No. US 1991-660162, filed on 22 Feb 1991 which is a continuation of Ser. No. US 1989-422699, filed on

17 Oct 1989 which is a continuation-in-part of Ser. No. US 1989-315342, filed on 23 Feb 1989, now patented, Pat. No. US 5011691 And a continuation-in-part of Ser. No. US 1990-569920, filed on 20 Aug 1990, now abandoned

which is a continuation-in-part of Ser. No. US

1989-422699, filed on 17 Oct 1989 which is a continuation-in-part of Ser. No. US 1990-483913, filed on 22 Feb 1990, now patented, Pat. No. US 5171574 which is a continuation-in-part of Ser. No. US 1989-422613, filed on 17 Oct 1989, now patented, Pat. No. US 4975526 which is a continuation-in-part of Ser. No. US 1989-315342, filed on 23 Feb 1989, now patented, Pat. No. US 5011691 And a continuation-in-part of Ser. No. US 1990-569920, filed on 20 Aug 1990, now abandoned which is a continuation-in-part of Ser. No. US 1989-422699, filed on 17 Oct 1989 which is a continuation-in-part of Ser. No. US 1990-483913, filed on 22 Feb 1990, now patented, Pat. No. US 5171574 which is a continuation-in-part of Ser. No. US 1989-422613, filed on 17 Oct 1989, now patented, Pat. No. US 4975526 which is a continuation-in-part of Ser. No. US 1989-315342, filed on 23 Feb 1989, now patented, Pat. No. US 5011691 And a continuation-in-part of Ser. No. US 1990-600024, filed on 18 Oct 1990, now abandoned which is a continuation-in-part of Ser. No. US 1990-569920, filed on 20 Aug 1990, now abandoned And a continuation-in-part of Ser. No. US 1990-599543, filed on 18 Oct 1990, now abandoned which is a continuation-in-part of Ser. No. US 1990-569920, filed on 20 Aug 1990, now abandoned And a continuation-in-part of Ser. No. US 1990-616374, filed on 21 Nov 1990, now patented, Pat. No. US 5162114 which is a division of Ser. No. US 1989-422613, filed on 17 Oct 1989, now patented, Pat. No. US 4975526 And a continuation-in-part of Ser. No. US 1990-483913, filed on 22 Feb 1990, now patented, Pat. No. US 5171574

DOCUMENT TYPE: Utility FILE SEGMENT: Granted

PRIMARY EXAMINER: Nutter, Nathan M.

LEGAL REPRESENTATIVE: Testa, Hurwitz & Thibeault

NUMBER OF CLAIMS: 58 EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 80 Drawing Figure(s); 47 Drawing Page(s)

LINE COUNT: 4211

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Disclosed are (1) osteogenic devices comprising a matrix containing substantially pure natural-sourced mammalian osteogenic protein; (2) DNA and amino acid sequences for novel polypeptide chains useful as subunits of dimeric osteogenic proteins; (3) vectors carrying sequences encoding these novel polypeptide chains and host cells transfected with these vectors; (4) methods of producing these polypeptide chains using recombinant DNA technology; (5) antibodies specific for these novel polypeptide chains; (6) osteogenic devices comprising these recombinantly produced proteins in association with an appropriate carrier matrix; and (7) methods of using the osteogenic devices to mimic the natural course of endochondral bone formation in mammals.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L17 ANSWER 36 OF 40 USPATFULL on STN

ACCESSION NUMBER: 94:35189 USPATFULL

TITLE: Prosthetic articular cartilage

INVENTOR(S): Stone, Kevin R., Mill Valley, CA, United States

Li, Shu-Tung, Oakland, NJ, United States

PATENT ASSIGNEE(S): ReGen Corporation, San Francisco, CA, United States

KIND

(U.S. corporation)

NUMBER

PATENT INFORMATION:	US 5306311	19940426	<
APPLICATION INFO.:	US 1991-809003	19911217	(7)
DISCLAIMER DATE:	20080416		

RELATED APPLN. INFO.: Continuation-in-part of Ser. No. US 1990-520027, filed on 19 May 1990 which is a continuation-in-part of Ser. No. US 1989-317951, filed on 2 Mar 1989, now patented, Pat. No. US 5007934 which is a continuation-in-part of

Ser. No. US 1987-75352, filed on 20 Jul 1987, now

DATE

patented, Pat. No. US 4880429

DOCUMENT TYPE: Utility FILE SEGMENT: Granted

PRIMARY EXAMINER: Isabella, David LEGAL REPRESENTATIVE: Lappin & Kusmer

NUMBER OF CLAIMS: 27 EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 12 Drawing Figure(s); 6 Drawing Page(s)

LINE COUNT: 955

AB A prosthetic, resorbable articular cartilage and methods of its fabrication and insertion are disclosed. The prosthetic articular cartilage, when implanted in a humanoid joint, acts as a resorbable scaffold for ingrowth of native articular chondrocytes and supports natural articulating joint forces. The prosthetic articular cartilage is a dry, porous, volume matrix of biocompatible and bioresorbable fibers. These fibers include a natural polymer or analogs thereof, at least a portion of which may be crosslinked. The matrix is adapted to have an in vivo outer surface contour substantially the same as that of natural articular cartilage in an articulating joint, and has a pore size in the approximate range of about 100 microns to about 400 microns.

L17 ANSWER 37 OF 40 USPATFULL on STN

ACCESSION NUMBER: 93:100856 USPATFULL TITLE: Osteogenic proteins

INVENTOR(S): Oppermann, Hermann, Medway, MA, United States Ozkaynak, Engin, Milford, MA, United States

Kuberasampath, Thangavel, Medway, MA, United States Rueger, David C., Hopkinton, MA, United States

Pang, Roy H. L., Medway, MA, United States
PATENT ASSIGNEE(S): Stryker Corporation, Kalamazoo, MI, United States (U.S.

corporation)

	NUMBER	KIND DAT	ΓE	
PATENT INFORMATION:	US 5266683	19931	1130	<
APPLICATION INFO.:	US 1992-841646	19920	0221 (7)	
DISCLAIMER DATE:	20101102			

RELATED APPLN. INFO.: Continuation-in-part of Ser. No. US 1992-827052, filed

on 28 Jan 1992, now patented, Pat. No. US 5250302 Ser. No. Ser. No. US 1990-579865, filed on 7 Oct 1990, now patented, Pat. No. US 5108753 Ser. No. Ser. No. US 1990-621849, filed on 4 Dec 1990, now abandoned Ser. No. Ser. No. US 1990-621988, filed on 4 Dec 1990, now abandoned Ser. No. Ser. No. US 1991-810560, filed on 20

Dec 1991, now abandoned Ser. No. Ser. No. US

1990-569920, filed on 20 Aug 1990, now abandoned Ser.

No. Ser. No. US 1990-600024, filed on 18 Oct 1990, now abandoned Ser. No. Ser. No. US 1990-599543, filed on 18 Oct 1990, now abandoned Ser. No. Ser. No. US 1990-616374, filed on 21 Nov 1990, now patented, Pat. No. US 5162114 And Ser. No. US 1990-483913, filed on 22 Feb 1990, now patented, Pat. No. US 5171574, said Ser. 827052 which is a division of Ser. No. US 1988-179406, filed on 8 Apr 1988, now patented, Pat. No. US 4968590 , said Ser. No. 579865 which is a 179406 , said Ser. No. division of Ser. No. 621849 which is a division of Ser. No. US 1988-232630, filed on 15 Aug 1988, now abandoned which is a continuation-in-part of Ser. No. 179406 , said Ser. 621988 which is a division of Ser. No. US No. 1989-315342, filed on 23 Feb 1989, now patented, Pat. No. US 5011691 which is a continuation-in-part of Ser. 810560 which is a 232630 , said Ser. No. continuation of Ser. No. US 1991-660162, filed on 22 Feb 1991, now abandoned which is a continuation of Ser. No. US 1989-422699, filed on 17 Oct 1989, now abandoned which is a continuation-in-part of Ser. No. said Ser. No. 569920 which is a continuation-in-part 422699 And Ser. No. 483913 which is a of Ser. No. continuation-in-part of Ser. No. US 1989-422613, filed on 17 Oct 1989, now patented, Pat. No. US 4975526 which is a continuation-in-part of Ser. No. 315342 , said Ser. No. 600024 which is a continuation-in-part of 569920 , said Ser. No. 599543 which is a Ser. No. 569920 which is a continuation-in-part of Ser. No. continuation-in-part of Ser. No. 569920

DOCUMENT TYPE: Utility FILE SEGMENT: Granted

PRIMARY EXAMINER: Nutter, Nathan M.

LEGAL REPRESENTATIVE: Testa, Hurwitz & Thibeault

NUMBER OF CLAIMS: 58
EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 76 Drawing Figure(s); 47 Drawing Page(s)

LINE COUNT: 4144

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Disclosed are (1) osteogenic devices comprising a matrix containing substantially pure natural-sourced mammalian osteogenic protein; (2) DNA and amino acid sequences for novel polypeptide chains useful as subunits of dimeric osteogenic proteins; (3) vectors carrying sequences encoding these novel polypeptide chains and host cells transfected with these vectors; (4) methods of producing these polypeptide chains using recombinant DNA technology; (5) antibodies specific for these novel polypeptide chains; (6) osteogenic devices comprising these recombinantly produced proteins in association with an appropriate carrier matrix; and (7) methods of using the osteogenic devices to mimic the natural course of endochondral bone formation in mammals.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L17 ANSWER 38 OF 40 USPATFULL on STN

ACCESSION NUMBER: 93:97929 USPATFULL TITLE: Prosthetic ligaments

INVENTOR(S): Li, Shu-Tung, Oakland, NJ, United States

Stone, Kevin R., Mill Valley, CA, United States

PATENT ASSIGNEE(S): ReGen Biologics, Inc., San Francisco, CA, United States

(U.S. corporation)

NUMBER KIND DATE

PATENT INFORMATION:	US 5263984 19931123 <
APPLICATION INFO.:	US 1992-872636 19920422 (7)
RELATED APPLN. INFO.:	Continuation-in-part of Ser. No. US 1990-582516, filed
THE STATE OF THE S	on 13 Sep 1990, now patented, Pat. No. US 5116374 which
	is a division of Ser. No. US 1989-317951, filed on 2
	Mar 1989, now patented, Pat. No. US 5007934, issued on
	16 Apr 1991 which is a continuation-in-part of Ser. No.
	US 1987-75352, filed on 20 Jul 1987, now patented, Pat.
	No. US 4880429, issued on 14 Nov 1989
DOCUMENT TYPE:	Utility
FILE SEGMENT:	Granted
PRIMARY EXAMINER:	Isabella, David
LEGAL REPRESENTATIVE:	Lahive & Cockfield
NUMBER OF CLAIMS:	21
EXEMPLARY CLAIM:	1
NUMBER OF DRAWINGS:	<pre>10 Drawing Figure(s); 4 Drawing Page(s)</pre>
LINE COUNT:	755
	prosthetic ligament comprising a Plurality of
	aligned, elongated filaments. Each filament is a dry,
	matrix of biocompatible and bioresorbable fibrils, at
	which are crosslinked. The fibrils are short segments of
	of polymeric connective tissue components, or analogs
	Filament establishes a bioresorbable scaffold adapted for
	gament fibroblasts, the scaffold and the
	lasts supporting natural ligament tensile forces.
	are methods of fabricating the prosthetic ligament, and
methods of rege	enerating ligamentous tissue in vivo.
L17 ANSWER 39 OF 40 ACCESSION NUMBER: TITLE: INVENTOR(S): PATENT ASSIGNEE(S):	84:66147 USPATFULL Bone-equivalent and method for preparation thereof Bell, Eugene, Dedham, MA, United States Massachusetts Institute of Technology, Cambridge, MA,
ACCESSION NUMBER: TITLE: INVENTOR(S):	84:66147 USPATFULL Bone-equivalent and method for preparation thereof Bell, Eugene, Dedham, MA, United States Massachusetts Institute of Technology, Cambridge, MA, United States (U.S. corporation) NUMBER KIND DATE
ACCESSION NUMBER: TITLE: INVENTOR(S): PATENT ASSIGNEE(S):	84:66147 USPATFULL Bone-equivalent and method for preparation thereof Bell, Eugene, Dedham, MA, United States Massachusetts Institute of Technology, Cambridge, MA, United States (U.S. corporation) NUMBER KIND DATE
ACCESSION NUMBER: TITLE: INVENTOR(S): PATENT ASSIGNEE(S): PATENT INFORMATION:	84:66147 USPATFULL Bone-equivalent and method for preparation thereof Bell, Eugene, Dedham, MA, United States Massachusetts Institute of Technology, Cambridge, MA, United States (U.S. corporation) NUMBER KIND DATE
ACCESSION NUMBER: TITLE: INVENTOR(S): PATENT ASSIGNEE(S): PATENT INFORMATION: APPLICATION INFO.:	84:66147 USPATFULL Bone-equivalent and method for preparation thereof Bell, Eugene, Dedham, MA, United States Massachusetts Institute of Technology, Cambridge, MA, United States (U.S. corporation) NUMBER KIND DATE
ACCESSION NUMBER: TITLE: INVENTOR(S): PATENT ASSIGNEE(S): PATENT INFORMATION:	84:66147 USPATFULL Bone-equivalent and method for preparation thereof Bell, Eugene, Dedham, MA, United States Massachusetts Institute of Technology, Cambridge, MA, United States (U.S. corporation) NUMBER KIND DATE US 4485097 19841127 < US 1983-497984 19830525 (6) Continuation-in-part of Ser. No. US 1982-381978, filed
ACCESSION NUMBER: TITLE: INVENTOR(S): PATENT ASSIGNEE(S): PATENT INFORMATION: APPLICATION INFO.:	84:66147 USPATFULL Bone-equivalent and method for preparation thereof Bell, Eugene, Dedham, MA, United States Massachusetts Institute of Technology, Cambridge, MA, United States (U.S. corporation) NUMBER KIND DATE
ACCESSION NUMBER: TITLE: INVENTOR(S): PATENT ASSIGNEE(S): PATENT INFORMATION: APPLICATION INFO.:	84:66147 USPATFULL Bone-equivalent and method for preparation thereof Bell, Eugene, Dedham, MA, United States Massachusetts Institute of Technology, Cambridge, MA, United States (U.S. corporation) NUMBER KIND DATE
ACCESSION NUMBER: TITLE: INVENTOR(S): PATENT ASSIGNEE(S): PATENT INFORMATION: APPLICATION INFO.:	84:66147 USPATFULL Bone-equivalent and method for preparation thereof Bell, Eugene, Dedham, MA, United States Massachusetts Institute of Technology, Cambridge, MA, United States (U.S. corporation) NUMBER KIND DATE
ACCESSION NUMBER: TITLE: INVENTOR(S): PATENT ASSIGNEE(S): PATENT INFORMATION: APPLICATION INFO.:	84:66147 USPATFULL Bone-equivalent and method for preparation thereof Bell, Eugene, Dedham, MA, United States Massachusetts Institute of Technology, Cambridge, MA, United States (U.S. corporation) NUMBER KIND DATE
ACCESSION NUMBER: TITLE: INVENTOR(S): PATENT ASSIGNEE(S): PATENT INFORMATION: APPLICATION INFO.:	84:66147 USPATFULL Bone-equivalent and method for preparation thereof Bell, Eugene, Dedham, MA, United States Massachusetts Institute of Technology, Cambridge, MA, United States (U.S. corporation) NUMBER KIND DATE
ACCESSION NUMBER: TITLE: INVENTOR(S): PATENT ASSIGNEE(S): PATENT INFORMATION: APPLICATION INFO.:	84:66147 USPATFULL Bone-equivalent and method for preparation thereof Bell, Eugene, Dedham, MA, United States Massachusetts Institute of Technology, Cambridge, MA, United States (U.S. corporation) NUMBER KIND DATE
ACCESSION NUMBER: TITLE: INVENTOR(S): PATENT ASSIGNEE(S): PATENT INFORMATION: APPLICATION INFO.:	84:66147 USPATFULL Bone-equivalent and method for preparation thereof Bell, Eugene, Dedham, MA, United States Massachusetts Institute of Technology, Cambridge, MA, United States (U.S. corporation) NUMBER KIND DATE
ACCESSION NUMBER: TITLE: INVENTOR(S): PATENT ASSIGNEE(S): PATENT INFORMATION: APPLICATION INFO.: RELATED APPLN. INFO.:	84:66147 USPATFULL Bone-equivalent and method for preparation thereof Bell, Eugene, Dedham, MA, United States Massachusetts Institute of Technology, Cambridge, MA, United States (U.S. corporation) NUMBER KIND DATE
ACCESSION NUMBER: TITLE: INVENTOR(S): PATENT ASSIGNEE(S): PATENT INFORMATION: APPLICATION INFO.: RELATED APPLN. INFO.:	84:66147 USPATFULL Bone-equivalent and method for preparation thereof Bell, Eugene, Dedham, MA, United States Massachusetts Institute of Technology, Cambridge, MA, United States (U.S. corporation) NUMBER KIND DATE
ACCESSION NUMBER: TITLE: INVENTOR(S): PATENT ASSIGNEE(S): PATENT INFORMATION: APPLICATION INFO.: RELATED APPLN. INFO.: DOCUMENT TYPE: FILE SEGMENT: PRIMARY EXAMINER:	84:66147 USPATFULL Bone-equivalent and method for preparation thereof Bell, Eugene, Dedham, MA, United States Massachusetts Institute of Technology, Cambridge, MA, United States (U.S. corporation) NUMBER KIND DATE
ACCESSION NUMBER: TITLE: INVENTOR(S): PATENT ASSIGNEE(S): PATENT INFORMATION: APPLICATION INFO.: RELATED APPLN. INFO.: DOCUMENT TYPE: FILE SEGMENT: PRIMARY EXAMINER: LEGAL REPRESENTATIVE:	84:66147 USPATFULL Bone-equivalent and method for preparation thereof Bell, Eugene, Dedham, MA, United States Massachusetts Institute of Technology, Cambridge, MA, United States (U.S. corporation) NUMBER KIND DATE
ACCESSION NUMBER: TITLE: INVENTOR(S): PATENT ASSIGNEE(S): PATENT INFORMATION: APPLICATION INFO.: RELATED APPLN. INFO.: DOCUMENT TYPE: FILE SEGMENT: PRIMARY EXAMINER: LEGAL REPRESENTATIVE: NUMBER OF CLAIMS:	84:66147 USPATFULL Bone-equivalent and method for preparation thereof Bell, Eugene, Dedham, MA, United States Massachusetts Institute of Technology, Cambridge, MA, United States (U.S. corporation) NUMBER KIND DATE
ACCESSION NUMBER: TITLE: INVENTOR(S): PATENT ASSIGNEE(S): PATENT INFORMATION: APPLICATION INFO.: RELATED APPLN. INFO.: DOCUMENT TYPE: FILE SEGMENT: PRIMARY EXAMINER: LEGAL REPRESENTATIVE: NUMBER OF CLAIMS: EXEMPLARY CLAIM:	84:66147 USPATFULL Bone-equivalent and method for preparation thereof Bell, Eugene, Dedham, MA, United States Massachusetts Institute of Technology, Cambridge, MA, United States (U.S. corporation) NUMBER KIND DATE
ACCESSION NUMBER: TITLE: INVENTOR(S): PATENT ASSIGNEE(S): PATENT INFORMATION: APPLICATION INFO.: RELATED APPLN. INFO.: DOCUMENT TYPE: FILE SEGMENT: PRIMARY EXAMINER: LEGAL REPRESENTATIVE: NUMBER OF CLAIMS: EXEMPLARY CLAIM: NUMBER OF DRAWINGS:	84:66147 USPATFULL Bone-equivalent and method for preparation thereof Bell, Eugene, Dedham, MA, United States Massachusetts Institute of Technology, Cambridge, MA, United States (U.S. corporation) NUMBER KIND DATE
ACCESSION NUMBER: TITLE: INVENTOR(S): PATENT ASSIGNEE(S): PATENT INFORMATION: APPLICATION INFO.: RELATED APPLN. INFO.: DOCUMENT TYPE: FILE SEGMENT: PRIMARY EXAMINER: LEGAL REPRESENTATIVE: NUMBER OF CLAIMS: EXEMPLARY CLAIM:	84:66147 USPATFULL Bone-equivalent and method for preparation thereof Bell, Eugene, Dedham, MA, United States Massachusetts Institute of Technology, Cambridge, MA, United States (U.S. corporation) NUMBER KIND DATE

AB A bone-equivalent, useful in the fabrication of prostheses, is disclosed

which is prepared from a hydrated collagen lattice contracted by fibroblast cells and containing demineralized bone powder.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L17 ANSWER 40 OF 40 USPATFULL on STN

84:66146 USPATFULL ACCESSION NUMBER:

Tissue-equivalent and method for preparation thereof TITLE:

Bell, Eugene, Dedham, MA, United States INVENTOR(S):

Massachusetts Institute of Technology, Cambridge, MA, PATENT ASSIGNEE(S):

United States (U.S. corporation)

KIND DATE NUMBER _____ US 4485096 19841127 PATENT INFORMATION:

US 1982-381978 <--19820526 (6)

APPLICATION INFO.: Continuation-in-part of Ser. No. US 1982-352586, filed RELATED APPLN. INFO.:

on 26 Feb 1982, now abandoned which is a

continuation-in-part of Ser. No. US 1981-245536, filed

on 19 Mar 1981, now abandoned which is a

continuation-in-part of Ser. No. US 1978-972832, filed

on 26 Dec 1978, now abandoned

DOCUMENT TYPE: Utility FILE SEGMENT: Granted PRIMARY EXAMINER: Rosen, Sam

LEGAL REPRESENTATIVE: Smith, Jr., Arthur A., Brook, David E.

NUMBER OF CLAIMS: 29 EXEMPLARY CLAIM: 1,10

14 Drawing Figure(s); 7 Drawing Page(s) NUMBER OF DRAWINGS:

LINE COUNT: 1054

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

A tissue-equivalent, useful in the treatment of burns or other skin wounds and in the fabrication of prostheses, is disclosed which is prepared from a hydrated collagen lattice contracted by a contractile agent, such as fibroblast cells or blood platelets, to form tissue-equivalent. In one embodiment, a skin-equivalent can be fabricated by growing a layer of keratinocyte cells thereon. Both the keratinocyte cells and contractile agent may be derived from the potential recipient of the skin-equivalent.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.